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Did Human Culture Emerge in a Cultural Evolutionary Transition in Individuality?

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Abstract

Evolutionary Transitions in Individuality (ETI) have been responsible for the major transitions in levels of selection and individuality in natural history, such as the origins of prokaryotic and eukaryotic cells, multicellular organisms, and eusocial insects. The integrated hierarchical organization of life thereby emerged as groups of individuals repeatedly evolved into new and more complex kinds of individuals. The Social Protocell Hypothesis (SPH) proposes that the integrated hierarchical organization of human culture can also be understood as the outcome of an ETI—one that produced a “cultural organism” (a “sociot”) from a substrate of socially learned traditions that were contained in growing and dividing social communities. The SPH predicts that a threshold degree of evolutionary individuality would have been achieved by 2.0–2.5 Mya, followed by an increasing degree of evolutionary individuality as the ETI unfolded. We here assess the SPH by applying a battery of criteria—developed to assess evolutionary individuality in biological units—to cultural units across the evolutionary history of *Homo*. We find an increasing agreement with these criteria, which buttresses the claim that an ETI occurred in the cultural realm.

Keywords Cultural evolution · Cultural group selection · Evolutionary transitions in individuality · Human evolution · Social protocell · Sociot

Introduction

Traditional behaviors are found in many animal species (Galef and Laland 2005; Allen 2019), but only in *Homo* have they coalesced into integrated and shared cultural systems (e.g., Andersson et al. 2014a; Smaldino 2014; Richerson et al. 2016; Buskell et al. 2019; Read and Andersson 2019), and only *Homo* has come to specialize in maintaining and acting within such systems (e.g., via expanded cognitive and metacognitive functions; Sherwood et al. 2008; Csibra and Gergely 2011; Whiten and Erdal 2012; Shea et al. 2014; Sherwood and Gómez-Robles 2017; Dunstone and Caldwell 2018). The integrated cultural nature of human behavior is strikingly expressed in emblematic feats of cooperation and coordination—such as when hunting like “a highly competitive group-level predator” (Whiten and Erdal 2012)—but it also permeates the human way of life entirely and into its most minute details. For example, resources obtained using cultural hunting and foraging strategies go on to enter an intricate cultural “metabolic system” where they are processed, stored, distributed, disposed, and turned into a wide variety of products. These products themselves are part of

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the operation of this cultural system, which is much wider and older than its individual human stewards, who depend on it for their survival (e.g., Boyd and Richerson 2000; Henrich and McElreath 2003). This uniquely human “cultural community” embodies an emergent ecological strategy that cannot be reduced either to its learned or its genetic components (1992).

In many ways, this sounds more like the description of an organism than of an animal social community—except perhaps for some species of social insects, whose communities can be understood as an unusual type of organism, due in part to the high degree of relatedness within colonies (Queller 2000; Queller and Strassmann 2009; Kennedy et al. 2017). In human cultural communities, however, these integrated qualities cannot be understood genetically. While clearly underpinned by genetic adaptations, the exceptional variation of behavior seen in human communities is not explained by genetic variation (e.g., Lewontin 1972; Foley and Lahr 2011). Our genetic adaptation in this regard is indirect—it *permits* cultural adaptation to happen.

So how could features seen as typical of (or even unique to) adapted biological organization emerge from a substrate of social learning? Are we just pushing a tempting analogy too far? The Social ProtoCell Hypothesis (SPH; Andersson and Törnberg 2019) proposes that this affinity between cultural and biological organization is genuine and fruitful, and that it stems from a deep similarity in how they originated and evolved. More specifically, the SPH proposes that groups of socially learned traditions were integrated into a new group-level cultural entity via an Evolutionary Transition in Individuality (ETI; see Maynard-Smith and Szathmáry 1995; Michod 1999, 2007; Leigh 2010; Clarke 2014; Hanschen et al. 2015; Szathmáry 2015). In this view, group-level cultural integration did not follow as a secondary effect of the evolution of lower-level factors that are usually seen as primary (e.g., cumulative traditions, hominin cooperation, etc.) but precisely the other way around.

ETIs are rare evolutionary transitions where new higher-level *evolutionary individuals* (entities equipped to undergo adaptation by natural selection as wholes; Lewontin 1970; Buss 1987; Sober and Wilson 1994) arise from cooperating groups of lower-level evolutionary individuals (Buss 1987; Szathmáry and Maynard-Smith 1995; Queller 2000; Michod and Roze 2001; Hanschen et al. 2018). Repeated ETIs have thereby produced one of life’s most familiar characteristics: its hierarchical structure. For example, groups of cooperative genes evolved into the first cellular genome, groups of bacteria-like cells evolved into the eukaryotic cell, groups of eukaryotic cells evolved into multicellular organisms, and groups of multicellular organisms evolved into social insect colonies.

Evolutionary individuality (Buss 1987; Michod 1999; Radzvilavicius and Blackstone 2018) emerges during an ETI

via cycles of cooperation, conflict, and conflict mediation. Selection and adaptation thereby move to a higher level of organization, producing an integrated type of entity whose lower-level units (originally evolutionary individuals) are co-opted and turned into parts of the whole (Michod 1999). For a brief overview of ETI theory and its relation to frameworks such as Major Transitions in Evolution, see Hanschen et al. (2018).

The cultural ETI described by the SPH is proposed to have started some 2.5 Mya, associated with the emergence of hominin big game carnivory and the appearance of the genus *Homo*. It is proposed to have been primed by a pre-existing fortuitous combination of behavioral and ecological circumstances that imparted a basic level of evolutionary individuality (via community-level boundaries, heredity, and reproduction) to collections of unintegrated traditions maintained in growing and splitting early hominin social communities.

The transition would have taken us from “animal culture” to integrated cultural entities, equipped with irreducible systems for heredity, reproduction, and development on the cultural group level. Traditions would evolve into increasingly subordinated components of a hierarchically organized cultural whole. Following Andersson and Törnberg (2019), the hypothetical type of evolutionary individual that emerged in this transition is referred to as a *sociont*. Notably, hominins are not seen as part of the emerging sociont but remain separate genetic evolutionary individuals.¹ The SPH thereby introduces a notable change of perspective since “the group” is here a group of traditions, not of hominins.

Arguably, the most fundamental question to pose about the SPH is whether—and, if so, when and to what degree—the culture of *Homo* shows evidence of evolutionary individuality. This question has many parts. For example, is there evidence that selection at an early point came to act collectively on combinations of animal-style traditions in hominin social communities? Were integrated systems of traditions formed as a result? Is the function and organization of *Homo* culture consistent with what one would expect if selection increasingly acted on cultural systems as wholes? Is there evidence that these proposed cultural organisms (like biological counterparts) evolved mechanisms that increased the extent to which they could be targets of selection? These are the questions we seek to address in this article.

We examine these issues by applying a set of criteria for assessing whether biological entities qualify as evolutionary

¹ We let the term “cultural community” represent the manifestation of what Andersson and Törnberg (2019) proposed as an obligate mutualism between *Homo* and sociont – i.e., the combined units of humans and culture. The nature of this relationship, and the evolution of *Homo* within it, is backgrounded in this article, which is focused on the emergence of the sociont as a cultural evolutionary individual.

individuals (Hanschen et al. 2018), adapting them to the cultural realm to account for the difference in substrate. We begin by outlining the SPH to explain why we think that an ETI drove human cultural evolution, and to introduce the entities used in the analysis. From this basis, we apply our criteria to judge whether, how, to what extent, and at roughly what stage they are fulfilled. The analyses are then summarized and compared with selected types of biological evolutionary individuals, and the results are assessed and compared with theoretical expectations. We conclude by discussing the results and evolutionary individuality in the context of a set of features of human culture that appear to be inconsistent with our results. This article combines schemata from several fields; to aid the reader we have provided a glossary for reference.

From Traditions to Socients via the Social Protocell: An Overview

Pan as a Proxy for Early Hominins

We use the traditions and community dynamics of *Pan* (in particular the more studied common chimpanzee *Pan troglodytes*) as a proxy for a primordial (pre-*Homo*) early hominin condition. Aware of the risk of conveniently overstating similarities between *Pan* and early hominins (e.g., Sayers and Lovejoy 2008), our arguments rest in particular on assumed similarities in the following basic aspects of group behavior, social learning, and ecological strategy.

The diverse and broad range of traditions maintained by *Pan* include extractive foraging behaviors such as nut-cracking, leaf sponging, termite fishing, and ant-dipping, along with a wide variety of social conventions, food choices, and so on (see Boesch 2012 for overview). These traditions are transmitted between individual apes primarily by copying outcomes (emulation) rather than underlying processes (imitation; Tomasello et al. 1987; Tomasello 1996; Tennie et al. 2009; Whiten et al. 2009; Clay and Tennie 2018), and they may be stable and potentially long-lived (Mercader et al. 2002, 2007). Extant chimpanzees are believed to be qualitatively similar to the earliest hominins with regard to their capacity to form and maintain traditions (e.g., Whiten et al. 2003; McGrew 2010; van Schaik 2016, p. 78), and it is likely that early hominins maintained traditions at a level and of a type similar to extant wild chimpanzees (e.g., Boesch and Tomasello 1998; Whiten et al. 1999, 2003; Whiten 2005; Lycett et al. 2009; Harmand et al. 2015).

With regard to overall community organization, Pleistocene *Homo* appears to evolve from the basis of a *Pan*-like fission–fusion type of organization, through increasing refinement and the addition of new and intermediate levels of social organization (Grove et al. 2012; Layton et al.

2012) —not by shifting to some radically different type of group organization. Of particular interest is the community lifecycle of *Pan* (Moffett 2013, pp. 239–249; Andersson and Törnberg 2019, p. 89). Communities arise and expire in irreversible, and roughly symmetric, fission events when social conflicts spiral out of control (Goodall 1986; Furuichi 1987; Feldblum et al. 2018). This becomes progressively more likely if group size increases and overburdens social cognitive mechanisms for handling conflicts and maintaining cohesion (Dunbar 1992, 1993, 1998). These events are under-researched but appear to be inherent to social features shared between *Pan* and *Homo*.

The Social Protocell

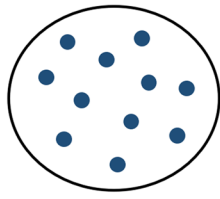
The centerpiece of the SPH is the so-called “social protocell” model, whose name derives from the protocell model of how early cells arose via an ETI in a substrate of primitive RNA molecules (Gánti 1975, 1997; Michod 1983; Szathmáry 1986; Szathmáry and Demeter 1987; Szathmáry and Maynard-Smith 1995; Norris and Raine 1998). The claim is that the evolution of the socient would have followed a similar pathway, but in a very different substrate. We will now briefly review the argument by Andersson and Törnberg (2019).

The social protocell is a set of circumstances that, as a side effect of their organization, creates the potential for selection to act on groups of traditions contained in social communities (see Fig. 1). This condition is claimed to be incompletely present in *Pan* communities today, and to thereby likely have existed also in early hominins (Andersson and Törnberg 2019, pp. 90–91; see also the previous section). The condition may be unpacked in terms of a system of three group-level evolutionary meta-functions: *boundaries*, *reproduction*, and *heredity* (see Table 1).

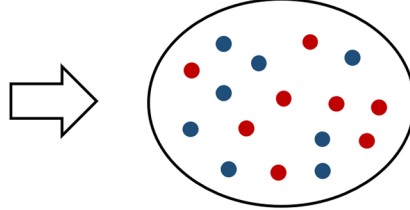
These functions potentiate the evolution of group-level *systems* of coadapted traditions that would substantially expand the range of solutions achievable by social learning. Quite simply, you can do more (and different) things with an emergent system of traditions than you can with single traditions. We here refer to such adapted integrated systems of cultural components as *institutional* (Richerson et al. 2016). The socient, in other words, consists of institutional organization. To illustrate, we outline “the Oldowan carnivory institution”² in Fig. 2 as a potential example of an early (ca. 2.6–1.8 Mya) institutional system of coadapted

² Oldowan technology was variable but saw a progression from simpler to more sophisticated strategies with regard to, for example, long-distance transportation of raw material, raw material selection, tool production, and hunting strategies. Our schematic illustration makes no claim to represent Oldowan culture as a whole or at any particular point in time or space.

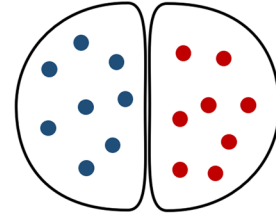
The **biological protocell** is a physical enclosure of proto-biotic RNA genes by a membrane vesicle generated as a metabolic by-product.



The growing vesicle becomes unstable and fissions spontaneously.



Daughter protocells divide the content of the parent, including the self-replicating RNA, creating two copies.



The **social protocell** is a socially generated enclosure of animal traditions. Social interactions are tight within but not between communities which causes containment of traditions since these spread across strong social links.

The social network becomes less resilient to conflict as the community grows. Eventually, two social sub-groups coalesce and the community fissions roughly symmetrically.

Dividing the community divides the social network, and if traditions spread rapidly inside communities, both these new networks will contain similar groups of traditions.

Fig. 1 The SPH proposes that social communities impose a group-level lifecycle on collections of traditions, in the same way that protocells did with regard to proto-biotic RNA genes. Above, we compare

idealized renditions of biological protocells with their proposed social counterparts to illustrate the parallelism

activities, each in distinct domains and contexts, and each likely supported by separately transmitted traditional behavior (e.g., Roche et al. 2009).

Fitness on the level of the social protocell would be driven by the biological fitness contributed by traditions to the hominins maintaining them. If the hominins survived and reproduced at higher rates, so would traditions contained within that social protocell (Fig. 1). But if traditions provided comparably small advantages, the fate of social protocells would be decided mostly by other factors, including the vagaries of chance. We therefore need reasons to infer that some important and widely available target, for which sophisticated institutional strategies would yield a substantial advantage, was available to early *Homo* but not to *Pan*, where an ETI was never initiated. Moreover, for evolution not to get stuck at an early point, this target must have kept yielding advantages as more and more sophisticated institutional strategies arose.

Cracking nuts or fishing for termites may provide adaptive additions to the diet, but they will hardly cause chimpanzee communities to decisively outcompete their neighbors. More generally, the rainforest resources available to *Pan* occur patchily and in small packages. Beyond a certain point of sophistication, the returns to increasing investments will thereby diminish. By contrast, *Homo* is uniquely associated with a resource that could have provided a strong and persistent competitive edge if pursued using cultural institutions, namely large carcasses. The earliest modified stone tools (the Oldowan complex) would have been especially useful for processing soft tissue on carcasses, and large game carnivory went on to become highly developed and foundational to the lifestyle of *Homo* during the Pleistocene—across a

widening variety of habitats and supported by sophisticated cultural systems (e.g., Stiner 2002; Bickerton and Szathmáry 2011; Whiten and Erdal 2012; Gintis et al. 2015).

But how would the social protocell get us from a *Pan*-like early hominin state to simple institutions? The australopithecine ancestors of *Homo* are believed to have gradually moved from facultative hunting of small animals (similar to *Pan*) toward obligate large game predation, presumably via facultative scavenging (for recent reviews see Thompson et al. 2019; Pobiner 2020). Although details are contested, most agree that this path involved ascending a gradient in risk and task complexity—driven by the benefits of accessing large packages of high-quality food, but facing a range of persistent hurdles, such as a need for processing and an increased exposure to predators and pathogens. Getting further in this direction would thereby put a premium on complex, coordinated, and cooperative strategies.³

As in *Pan* today, the social protocell would have been there all along as a side effect of component behaviors, maintained because they were adaptive in their own separate ways. Its incidental effect of enabling cultural group selection would kick in gradually as it enabled evolution in directions that otherwise could not be taken. The earliest stage could, for example, have

³ Hominin cooperation would obviously have been strongly enabling in just about any sociocultural institutional adaptation. Culturally mediated facilitation, manipulation, or punishment of individuals to enhance cooperation and coordination would thereby be an important aspect of sociocultural adaptation. Likewise, genetic adaptation as a cooperator in a cultural community would be an expected outcome of a mutualistic relation between *Homo* and socioculture (Andersson and Törnberg 2019, pp. 92–93).

Table 1 Mechanisms behind primitive evolutionary individuality in biological and social protocells

Feature	Biological Protocell	Social Protocell	Meta-evolutionary Function
<i>Higher-level boundaries</i>	RNA genes cannot pass through the lipid membrane vesicles that are generated as a by-product of their metabolism. They are thereby contained	Traditions spread easily across tight social networks within social communities, but only rarely between communities since social links are absent or weak	The boundaries concentrate and align the evolutionary fates of the contained units. This links their welfare partly to the welfare of their common container
<i>Higher-level reproduction</i>	As the vesicle grows by internal addition of more lipid molecules, it destabilizes and splits into two daughter vesicles	Communities tend to split roughly symmetrically at a rate that increases sharply as group size exceeds a critical threshold	Reproduction is necessary for higher-level fitness since differences between groups must result in differential success in competition with other containers
<i>Higher-level heredity</i>	When the vesicle splits into two daughter vesicles (subject to assumptions about size and symmetry) both will tend to contain the same mixes of RNA genes	When communities split, the populations of both daughter communities are likely to carry similar mixes of traditions, causing their overall traditional behavior to be similar	Heredity enables natural selection on the higher level since features that affect the fitness of the containers may persist, vary, and produce an evolutionary population

selectively preserved institutional elaborations on *Pan*-like traditional strategies, which perhaps involved late access to carcasses, without opposition, and the use of only immediately available materials such as unmodified rocks and bones. The social protocell would in this way have opened up an otherwise inaccessible cultural design space (see Stankiewicz 2000) that could keep expanding in co-evolution with its underpinning capacities in *Homo*.

The SPH proposes that the social protocell would thereby not simply boost cultural evolution but actually trigger the formation of the first-ever non-biotic unit of selection via an ETI. Key to understanding why we think an ETI may have unfolded is that the fortuitous meta-evolutionary functions imparted by the social protocell—i.e., boundaries, reproduction, and heredity (Table 1)—*themselves* could come under cultural control and be adaptively expanded and refined by the cultural group selection that they enabled. This can be understood theoretically as *the evolution of evolutionary individuality*.

Applying Evolutionary Individuality Criteria to Human Culture

Background

A variety of criteria have been proposed by researchers to define and test whether candidate entities qualify as evolutionary individuals. In the following sections we will use the most commonly applied criteria as reviewed by Hanschen et al. (2017): *spatial boundaries*, *informational uniqueness*, *informational homogeneity*, *indivisibility*, *group-level adaptations*, *division-of-labor*, and the applicability of a specific kind of multilevel selection termed *multilevel selection 2*. These criteria identify features that are generated by and/or enabling of group-level selection, and that are thereby likely to arise during an ETI, but unlikely to be seen otherwise (in particular together and in a highly developed state). In this way we aim to test the SPH and systematically articulate the hypothesis in an empirical context.

For each criterion we first explain its role and importance with regard to evolutionary individuality. We then interpret these criteria in terms of cultural systems in *Homo*. We emphasize the Plio-Pleistocene origin of the sociant via the social protocell, its early evolutionary history (primarily in the Oldowan), and, finally, we consider trends across the evolution of *Homo* during the Pleistocene.

Spatial/Temporal Boundaries

Description

Boundaries constrain the components of lower-level units in ETIs, keeping them from diffusing between groups and from

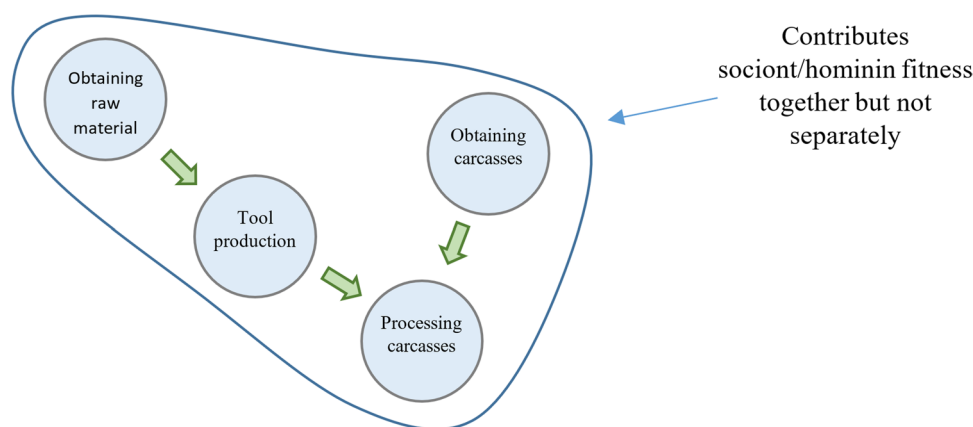


Fig. 2 Consider a minimal rendition of “the Oldowan carnivory institution.” The components are behavioral traditions that occupy distinctly different regimes in terms of time, location, type of behavior, and materials used. Traditions such as obtaining raw material, tool

production, and carcass processing are either pointless or impossible considered separately since they are adapted to be parts of an emergent system that, in turn, produces something useful for the sociant (via the hominins) as a whole

pursuing independent agendas that require free movement. For example, during the origin of cellular life, the protocellular lipid membrane kept autocatalytic chemical networks (the lower-level units in that transition) contained inside self-replicating vesicles (see Fig. 1). Being stuck together in this manner facilitated the evolution of cooperation and eventually the integration of dispersed genetic information into a genome (Jablonka and Szathmáry 1995; Maynard-Smith and Szathmáry 1995; Durand and Michod 2010); see also “the boomerang effect” (e.g., Dugatkin 2002).

Analysis

In “The Social Protocell” section, above, we claimed that collections of traditions were stuck together within the social protocell. The boundary in this case is a lack of social bridges across which transmission can happen between communities (Table 1). The social protocell and sociant are thereby primarily bounded in a social rather than a physical space, although for *Pan*, and frequently (but not always) also for *Homo*, this social space corresponds to a physical space in the form of a territory.

Three factors that cause robust and persistent containment of traditions in *Pan* are:

1. Close and persistent social contact favors the transmission of traditions. Such contact is present within communities but rarely applies between communities (e.g., Goodall 1986; Wilson and Wrangham 2003; Boesch et al. 2008; Schel et al. 2013).
2. Enculturated individuals cannot transfer freely between communities (e.g., Nishida et al. 1979; Pusey 1979; Wrangham 1979; Wilson and Wrangham 2003).

3. Enculturated individuals that do transfer are poor vectors of traditions, for example, due to conformity bias (Whiten et al. 2005; Van De Waal et al. 2010; Haun et al. 2012; Luncz and Boesch 2014, 2015) and rank bias (Horner et al. 2010; Kendal et al. 2015; Watson et al. 2017).

Although positive evidence is unavailable, it is quite plausible that the factors listed above would have applied similarly to early hominins in a Pliocene primordial state. These are primitive examples of what Durham (1992) labels *transmission isolating mechanisms*, which were subsequently expanded and institutionalized during the evolution of *Homo*.

One reason to suspect that the containment of culture increased rather than decreased in *Homo* is that the more complex culture became, the more strongly its transmission must have relied on close and persistent social intimacy. On the level of social learning, cultural components tend to be opaque (indeed, often both to role model and learner; e.g., Tostevin 2007, 2019; Premo and Tostevin 2016) and their transmission reliant on specialized “pedagogical” adaptations (Gergely and Csibra 2006; Tehrani and Riede 2008; Csibra and Gergely 2009, 2011; Kline 2015; Gärdenfors and Högberg 2017; Laland 2017). On higher levels of organization, increasingly complex and integrated hominin institutions may have diffused less and less easily (Richerson et al. 2016, p. 5): First, to be functional, all essential components of an institution must be effectively transmitted. Second, the institution, in turn, will be integrated into some specific higher-order system of institutions, which means it will likely be much less adaptive elsewhere. Third, institutions, even more than focused skills, rely on liberal amounts

of tacit knowledge whose function and even existence is unknown to the agents (Polanyi 1967).⁴

Direct empirical evidence of the timing and details are hard to come by, but interdisciplinary analysis suggests that the evolution of intercommunity boundaries has a gradual and drawn-out history across the Pleistocene (Grove et al. 2012). Layton et al. (2012) analyze data on the displacement of stone used for making artifacts (from Féblot-Augustins 1997) from the Oldowan to the Upper Paleolithic, in conjunction with estimated community sizes (Dunbar 1993; Hill and Dunbar 2003), and ethnographic as well as modeling analyses of area use. They conclude that movements of stone raw material remained mainly within small face-to-face coordinated social units (congruent with *Pan* communities) at least through the late Middle Paleolithic (until circa 50 kya, although simple intercommunity institutions may have emerged locally before that; e.g., Blegen 2017; Brooks et al. 2018). Exchange of lithic material may be expected to follow networks of amicable social interactions, and the spread of lithic material should therefore overlap with the transmission of culture. In late Pleistocene and Holocene human societies, social boundaries are certainly under cultural control and exhibit complex specializations (e.g., via cultural kinship, marriage, mythology, etc.; see, e.g., Read 2012). Not least language (e.g., via dialects) is a powerful boundary mechanism, and its function as such may have been an important factor in its evolution; see Moffett (2013, pp. 229–232).

Temporal boundaries are imposed by the irreversible community-level splitting dynamic, which the SPH views as analogous to cell division (see Fig. 1). Social protocells, and later socionts, thereby have beginnings and ends.

Summary

Boundaries in a social space here play the role that physical boundaries play in biology, and, although social spaces are frequently associated with physical territories, that is not always the case. These boundaries exist in *Pan* and thereby plausibly in early hominins, and rather than disappearing over time, they seem to have become more and more effective as barriers to culture, more institutionalized, and more subject to cultural adaptation.

⁴ Polanyi (1967) describes how tacit knowledge – i.e., knowledge we are unaware that we possess – hinders even simple acts of intentional transfer, even between highly similar contexts (e.g., moving equipment from one factory plant to another), even if done systematically, with high motivation, and large resources committed.

Informational Uniqueness

Description

In the biological realm, informational uniqueness essentially means that each unit has its own independent genetic makeup. Such units may exhibit heritable individual differences, which promote evolutionary individuality by enabling group-level variation that selection can act upon. In the SPH case, the heritable information is cultural rather than genetic, and we see the sociont as informationally unique to the extent that it possesses its own stable, independent, and heritable set of traditions or cultural components.

Analysis

Henrich (2004) describes and reviews evidence for four mechanisms that promote what we here call informational uniqueness; see also Boyd and Richerson (2010) and Chudek and Henrich (2011). These mechanisms suppress within-group variability and increase between-group variability in human behavior, assuming the presence of boundaries between groups. The first is *conformist* learning, which increases the rate of horizontal spread of favored traditions within a community and prevents established traditions from dropping out. The second is *prestige-biased* learning (Henrich and Gil-White 2001; Jiménez and Mesoudi 2019), which further increases between-group variation (e.g., Boyd and Richerson 1987) since it selectively disfavors ideas originating outside of the group, while it permits some internal sources of variation that can break up conformist lock-ins. The third is *punishment*; i.e., that the biological agents accept the cost of punishing nonconformers, which greatly amplifies the stabilizing effect of conformism (e.g., Boyd and Richerson 1992). Finally, *normative conformity* represents conformity for purely social reasons, regardless of whether the behavior in question is otherwise useful or not.

Conformism, *sensu lato*, is widespread among animals with social learning (de Waal 2013) and may contribute to maintaining between-group variation in chimpanzees (Whiten et al. 2005; van de Waal et al. 2010, 2013; Haun et al. 2012; van Leeuwen et al. 2012; Luncz and Boesch 2014, 2015). However, it is a weaker force in *Pan* than it is in humans, who conform not only to gain access to better information, but also normatively in pursuit of social benefits (e.g., Van Leeuwen et al. 2013; Haun et al. 2014). We thereby deem it likely that some form of conformism was present to some degree in early hominin communities and increased in *Homo* across the Pleistocene.

Chimpanzees may exhibit a bias towards learning from individuals with a high rank and/or a track record of success (Horner et al. 2010; Kendal et al. 2015; Watson et al. 2017). If present in early hominins, this may have worked

as an innate evolutionary starting point for “prestige” as a derived and culturally institutionalized version, buttressed by genetic adaptations (Henrich and Gil-White 2001). The other mechanisms reviewed by Henrich (2004) are weak, different, or absent in chimpanzees, and should consequently be viewed as derived during the evolution of *Homo*.

We argued above that *spatial/temporal boundaries* likely constrained horizontal (between-group) transmission before and (increasingly) during the evolution of *Homo*. Such boundaries to cultural dispersal, along with community-level heredity of *sets* of cultural components (via community-level splits; see “The Social Protocell” section), and the factors promoting informational uniqueness described above, indicate that communities may have diverged culturally over time due to selection or drift, rather than converging due to information flow—at least if cultural inheritance was sufficiently faithful.

Fidelity is important for the stable maintenance of informational uniqueness over time since it essentially bounds the amount of information that can be maintained by selection in a population (Eigen and Schuster 1977; Shea 2009; Andersson 2011, 2013). Notably, the fidelity of social protocell inheritance does not reduce to the fidelity of social learning, which may have been very low in early *Homo* (e.g., Tennie et al. 2017). Social protocell inheritance is simply the continuity of systems of specialized traditions through sociointer-reproduction events (see “The Social Protocell” section and Fig. 1). For example, say the abilities underpinning the Oldowan carnivory institution were, in some instance, distributed somewhat unevenly across a hundred hominins in a community. Fidelity may then be understood as the likelihood that, upon division, a sufficient number of instances of each needed tradition made it into the daughter communities to cause them to also feature this institution. This likelihood may be high also if the processes behind the traditions are transmitted with very low fidelity—as long as *their functions* are stable, which appears to potentially be the case (“*Pan as a Proxy for Early Hominins*” section).

Some degree of group-level informational uniqueness may thereby be theoretically expected in an early hominin *Pan*-like state. There is also some support for this in field studies where substantial between-group variation in traditions has been found between chimpanzee communities (e.g., Whiten et al. 1999, 2001; Sanz and Morgan 2007; Schöning et al. 2008; Lycett et al. 2009; Boesch 2012; Koops et al. 2015; van de Waal 2018; Kaufhold and Van Leeuwen 2019; Kalan et al. 2020).

Direct verification of informational uniqueness in Pleistocene *Homo* is challenging to obtain. Evidence is poorly synthesized and a coherent picture is lacking (Kuhn 2020). Numerous individual studies, however, support an overall picture of ancient and persistent geographical cultural heterogeneity. Analysis of traces of butchering techniques

at Bolomor Cave and Gran Dolina (Middle Pleistocene) shows evidence of persistent group-specific patterns that vary across time and space in ways that are not obviously functionally relevant (Blasco et al. 2013). Intercommunity technological variation has also been inferred archaeologically in the earliest Oldowan (at Gona; see Stout et al. 2010, 2019), and Chinese stone tool industries between 300 and 40 kya yield evidence of persistent regional cultural distinctiveness, despite their apparent simplicity (Bar-Yosef and Wang 2012; Gao 2013). Foley and Lahr (2011) moreover suggest that cultural transmission by expansion of groups best explains observed patterns of geographic cultural variation over the past 100,000 years.

Summary

Taken together, informational uniqueness on the level of communities is present to some extent in *Pan*, and clearly present in more recent *Homo*. It is also indirectly suggested to have been present, and to have increased, during the evolution of *Homo*.

Informational Homogeneity

Description

Informational homogeneity is maximized when all lower-level units in a biological individual carry the same genetic information. The fitness interests of the lower-level units are then aligned by the fact (and to the extent) that it makes no evolutionary difference which unit reproduces (Hamilton 1964a, b). Informational homogeneity thereby promotes group selection and provides an ideal setting for cooperation and so-called “fraternal” ETIs (Queller 1997) on the basis of kinship selection (e.g., Maynard-Smith and Szathmáry 1995; Michod 1999), such as when all cells in a clonally developing multicellular organism are genetically identical. In contrast, a low degree of homogeneity promotes competition and selection at the lower level, which can be problematic for the emergence of evolutionary individuality at the higher level.

Analysis

While informational homogeneity plays a central role in fraternal ETIs, an evolutionary individual that arises via an “egalitarian” ETI (Queller 1997; such as early cells arising from different species of RNA genes, or the eukaryotic cell arising from bacteria and archaeans) is *inherently* informationally heterogeneous. This egalitarian type of ETI would also be the best match for the SPH since different types of traditions and cultural components, specialized in different tasks, are clearly underpinned by different sets of

information. Components of culture are thereby more analogous to specialized types of genes becoming integrated into a genome than they are to clonal cells becoming integrated into a multicellular organism.

In our case, the informational homogeneity criterion is therefore neither expected to be met from the outset, nor to emerge as an outcome of the ETI. But the criterion is still important since the analysis tells us that lower-level competition will remain a problem if the scope of group selection is to keep expanding via the evolution of evolutionary individuality. During an egalitarian ETI, conflicts are resolved by the evolution of conflict modifier mechanisms (Michod and Nedelcu 2003). We should expect to find examples of this type of mechanisms in a sociont, as we do in cells.

If competition between traditions in the emerging sociont is analogous to competition between genes in the emerging cell, then cellular mechanisms for managing and suppressing genetic conflict may offer guidance. The integration of independently replicating RNA replicator/interactors into a genome is arguably the most central evolutionary innovation in this regard (Maynard-Smith and Szathmáry 1993). The chromosome is a specialized monopolistic group-level replicator whose operation is based on, but not reducible to, the original lower-level replication mechanism of independent genes. It replicates its genetic units, and thereby the group-level genetic structure and proportions, in a centralized and controlled manner once every lifecycle (Jablonka and Szathmáry 1995; Maynard-Smith and Szathmáry 1995; Durand and Michod 2010; Ågren 2014). This produces a setting where parasitic genes are suppressed since they need to become part of the chromosome to gain access to replication.

The evolution of the chromosome may exhibit a suggestive parallelism with the highly structured and institutionalized enculturation process that emerged in *Homo* (e.g., Read and Andersson 2019, pp. 2–3); see also the discussion of related processes by Smaldino (2014, pp. 250–251). A normative canon of cultural knowledge is here transmitted in a structured and cumulative sequence, following a modified and expanded process of physiological development (childhood), using adaptations for cultural transmission, that are unique to *Homo* (e.g., Thompson and Nelson 2011, 2016; Han and Ma 2015). Such an integration and monopolization of cultural heredity would stabilize higher-level cultural organization and make it more heritable, but it would also cause parasitic elements to be less likely to spread and disrupt the function of a sociont. The ability of cultural components to reproduce would become tied to admission into such a canon, which may require, for example, fitting functionally and logically into the prevailing system of customs and skills, and, not least, being considered part of the norm (see, e.g., contributions in Roughley and Bayertz 2019). Without such a normative centralized system, new traditions could

suddenly arise to exploit some feature of the sociont or the hominins (cognitively or psychologically) to spread and disrupt the integrated function of the sociont.

Summary

The sociont does not exhibit informational homogeneity, and it is not predicted by the SPH to do so. Provisionally an example of an egalitarian ETI, it should be expected to instead exhibit derived adaptations for suppressing lower-level competition between cultural components.

Indivisibility

Description

Indivisibility means that one cannot separate the parts out from the whole and maintain the functional properties of the whole. This increases the likelihood that selection will act on the integrated unit rather than on separate parts. It is therefore indicative of evolutionary individuality if separated subunits do not maintain properties of the whole and cannot survive on their own outside the group context (Michod 1999, 2007).

One mechanism by which indivisibility can emerge is when components specialize and lose vital features that are taken over by other specialized parts; see also the “[Evolutionary Division of Labor](#)” section, below. For example, cells in differentiated multicellular organisms have specialized in varied internal functions in the organism, and lost the ability to reproduce and survive independently in this process. The same fate has befallen bacterial mitochondria and plastid endosymbionts of eukaryotic cells, and, likewise, the specialized castes of social insects. Once this has happened, the fitness of one component is dependent upon other components of the group. Indivisibility indicates a low level of conflict between lower-level units since the dependencies that make the individual indivisible act to align fitness interests on the lower level.

Analysis

Since the sociont is composed of interacting cultural components, indivisibility means that components on any level of organization are unlikely to function well outside of the group context; see also the “[Informational Uniqueness](#)” section, where some problems pertaining to intercommunity transmission of cultural subunits were discussed. As a limit case, we may readily establish that institutions in modern-day human societies make little sense on their own. We could not take the institutions of a society and create functioning societies each using subsets of the parts—banking in one, police in the other, daycare in the first, and so on. There has to be a relatively

full set of complementary functional units in place. This logic permeates the entire internal hierarchy. Dividing institutional units on any level in this way incurs exactly the same set of problems, and the principle is particularly clearly expressed in technological systems, which are eminently indivisible.

This form of indivisibility is inherent to modular adapted systems, which we argued may have emerged early (circa 2.6–1.8 Mya); see the “Oldowan carnivory institution” (“[The Social Protocell](#)” section and Fig. 2). By contrast, sets of traditions contained in *Pan* communities are divisible *in principle*. Removing or adding one traditional practice—such as nut cracking or ant dipping—is unlikely to affect the function or transmission of other traditions since they lack interdependencies. The same would be true for a collection of early RNA species compartmentalized by a lipid membrane boundary. However, due to the protocellular dynamics (see Fig. 1 and Table 1) they would still not be divided regularly *in practice*. They would typically remain together, which would favor the evolution of dependencies, and thereby *actual* indivisibility.

Finally, while the division of sociants happens via the division of social groups of hominins, the former should not be confused with the latter (see “[The Social Protocell](#)” section and Fig. 2). Strictly speaking, the sociant never really divides (see “[Analysis](#)” of the “[Informational Uniqueness](#)” section). It simply persists in the daughter communities after a split if hominins carrying all necessary traditions make it across. Notably, this operation is less straightforward in the presence of division-of-labor between hominins (craft specialization). Cliques of hominins would then carry different parts of the cultural information, and additional mechanisms would be needed to ensure that hominins of all specializations make it into both daughter communities. Craft specialization is a feature of sedentary cultural communities, and despite good reasons for believing it to be generally adaptive on a group level (e.g., Henrich and Boyd 2008), it emerged only very late in prehistory.

Summary

Sets of animal traditions contained in the social protocell are divisible in principle but rarely divided in practice. Actual indivisibility would then have arisen during the evolution of the sociant, with the sociant becoming increasingly indivisible the more complex its institutional organization became.

Group-Level Adaptations

Description

Group-level adaptations provide evidence of group selection but identifying them can be challenging. Groups can have features that look like group-level adaptations but

that really are properties driven by selection on the lower level that filter up to the level of a group (Shelton and Michod 2014, 2020). Williams (1966) illustrated this by way of describing how a “fleet herd of deer” is really just a “herd of fleet deer” where the group-level property may be described as a “fortuitous benefit” (Williams 1966) or a “cross-level byproduct” (Okasha 2006) of lower-level properties.

Key to telling true group-level adaptations from cross-level byproducts is to determine whether fitness has truly been “exported” from the lower level to the group level, or if the fitness of the group is simply an aggregative property of lower-level traits (Michod and Herron 2006; Michod 2007). In other words, have the lower-level units sacrificed their fitness as independent individuals in return for a greater contribution of fitness via the higher level? We may subject claims of group-level adaptation to a test by asking whether carrying implicated traits would cause the lower-level entities to suffer a reduction of fitness if they left the context of the group. Being fleet, for example, fails this test since being fleet would not be detrimental to a deer if it left the group.

Analysis

Do the components of cultural systems have properties that would cause them to have lower fitness if they left their cultural context? Are such properties linked to adaptive properties of the cultural system that they benefit from being part of? If so, we may be looking at cultural traits selected on the group level.

Richerson et al. (Richerson et al. 2016) conclude that institutions are group-level features; see also Smaldino (2014). Although their analysis is mostly set in relatively recent times, we think a similar argument can be made from very early on. If we pose the question formulated above about the Oldowan carnivory institution (Fig. 2), we find that its constituent components will certainly suffer if moved to another setting, and that they will do so *because of* how they are adapted to serve their roles in the institution as an integrated whole. For example, making stone tools without the knowledge of how to obtain animal carcasses would be minimally beneficial and perhaps maladaptive. This contrasts with animal traditions (such as nut cracking or termite fishing among chimpanzees) which would seem to be equally adaptive regardless of the context of other traditions.⁵ Institutions then become more and more prevalent and complex the closer we get to modern times.

⁵ It should be noted that group-level features in primates is an under-researched topic and that it cannot be ruled out that marginal examples would exist (see, e.g., van de Waal 2018).

Summary

Group-level adaptations are absent (or marginal and not so far detected) in *Pan* communities. They seem to have arisen early during the evolution of *Homo*, whereafter they increased in complexity, integration, and importance. Institutions and institutions such as large game hunting may be early group-level traits.

Evolutionary Division of Labor

Description

To avoid confusion, let us first state that division of labor in anthropology (and in social science generally) refers specifically to craft specialization; i.e., the division of tasks and specializations between human individuals (e.g., Kuhn and Stiner 2006). Evolutionary theory uses a more general understanding of division of labor as a division of tasks and specializations between any types of components. For example, differentiated multicellular organisms exhibit division of labor between specialized cell types, cells exhibit division of labor between organelles, and social insect colonies exhibit division of labor between castes. Since cultural components rather than hominins form the parts of the sociont, we here refer to division of labor as occurring between cultural components.

Evidence of division of labor in an entity is evidence of “near-decomposability,” which is a universal principle of organization and design (see, e.g., Simon 1962; Wimsatt 1975; Marengo and Dosi 2005; Andersson and Törnberg 2018, pp. 129–31) where specialized functions are broken down into a level hierarchy of complementary sub-functions. This modular organization greatly simplifies and structures internal organization, and is evident in all but the very simplest adapted entities. It is simultaneously an outcome and a precondition for the evolution or design of complex adaptive organization. In an evolutionary context, division of labor is thereby evidence that fitness has been exported to the level of the group, and that the new higher-level entity has gained substantial evolutionary individuality.

Analysis

Archaeology robustly reveals a trend toward deepening division of labor between components of culture, both as observed in the products of culture (e.g., complex technology; see Querbes et al. 2014; Haidle et al. 2015), and in what we refer to as institutions; see also Smaldino (2014). Under the criteria of *Indivisibility* and *Group-level adaptations* we have already described several examples of the latter, and

the principle is clearly manifested in the Oldowan carnivory institution (Fig. 2), whose components are specialized in focused sub-tasks, most of which make sense only together with the other coadapted components.

This trend of diversification, hierarchization, and increasing narrowness of specialization of cultural components has continued and accelerated into the present (e.g., expressed as diversity of cultural products; see Beinhocker 2006). *Homo* is qualitatively different from other animals, including *Pan*, in this regard. The traditions maintained by chimpanzees exhibit diversification but *not* integration of function beyond what can be achieved cognitively in creative problem solving (within the “zone of latent solutions”; e.g., Tennie et al. 2009; Reindl et al. 2018). We take this to imply that pre-Oldowan hominin traditional repertoires (like *Pan*) likely did not exhibit division of labor.

Notably, as the sociont is the outcome of an egalitarian ETI (see “Analysis” of the “Informational Homogeneity” section),⁶ the pattern of division-of-labor in the sociont should more resemble what we see in unicellular organisms than what we see in multicellular organisms. Without informational homogeneity (see above) there is, for example, no basis for the differentiation between reproductive and somatic components that plays a key role in fraternal ETIs (e.g., single- to multicellular) by strongly suppressing lower-level selection.

Summary

Our observations lead us to conclude that cultural evolutionary division of labor arose early and deepened during the evolution of *Homo*, and that it is not evident in other species maintaining traditions, including *Pan*.

Multilevel Selection 2

Description

Damuth and Heisler (1988) seminally described a subdivision in the debate about multilevel selection in terms of two types of models: Multilevel Selection 1 (MLS1) and Multilevel Selection 2 (MLS2.) They characterized these as follows (“individual” here corresponds to our use of the term “lower-level”):

The criteria for MLS1 are as follows:

1. “Group selection” refers to the effects of group membership on individual fitness.

⁶ With traditions on the lower level – not a fraternal ETI with hominins on the lower level.

2. Fitnesses are properties of individuals and group fitness is an aggregative property of individual fitnesses.
3. Characters are values attributed to individuals (including both individual and contextual characters—see below).
4. Populations consist of individuals, organized into groups.
5. Explicit inferences can be made only about the changing proportions of different kinds of individuals in the whole population (the meta-population).

The criteria for MLS2 are as follows:

1. "Group selection" refers to change in the frequencies of different kinds of groups.
2. Fitnesses are properties of groups.
3. Characters are values attributed to groups (including both aggregate and global characters).
4. Populations consist of groups, composed of individuals.
5. Explicit inferences can be made only about the changing proportions of different kinds of groups in the population.

In essence, MLS2 models correspond to groups that are well on their way to becoming evolutionary individuals (Okasha 2006), while MLS1 models correspond to groups whose members do not constitute parts of a group-level individual, and where we cannot speak of group-level adaptations. Okasha (2007) furthermore remarked that there is a characteristic temporal ordering where MLS1 may turn into MLS2, so both types of dynamics may be in place at the same time.

Analysis

The ETI proposed by the SPH begins from an MLS1 scenario in which traditions are not integrated as parts of a higher-level system, but simply happen to be organized into groups as a by-product of early hominin group behavior (see "The Social Protocell" section). The fitness of such a group is an aggregative property of the effects of the different traditions since any interactions between them are minimal and likely not synergistic. As a result of these minimal interactions, characters and fitnesses may be assigned to individual traditions⁷ but not to groups thereof, except in the simple aggregate sense.

As the ETI progresses, institutional organization is argued to appear, as exemplified by the Oldowan carnivory institution (Fig. 2). Traditions here become specialized parts of

larger functional systems. The adaptive functions of such institutional structure (such as contributing meat or other resources) are emergent, as they are determined by complex, nonlinear interactions between its traditional components. We may thereby speak increasingly of fitness on the cultural group level rather than on the cultural component levels, and we are also more and more inclined to speak also of properties of these systems as wholes. Since traditions interact and co-occur, change on the levels of institutions and socients can also be characterized across time and space. In other words, traditions start out as independent entities, but they end up as specialized parts of larger systems.

Summary

The observations that we have made in prior sections support an interpretation where an MLS1 situation gradually turns into an MLS2 situation during the evolution of *Homo*. We deem that that the socient is consistent with a MLS2 framework, although, as in other egalitarian ETI, selection on the original level is never fully eliminated.

Results

In Table 2 we summarize our findings along with biological examples for comparison. The **eukaryotic cell** (like the socient) stems from an egalitarian ETI (eukaryogenesis) and is also the starting point of a fraternal ETI, namely the evolution of **multicellular organisms**. **Colonial organisms that develop clonally** (e.g., the volvocine green algae *Eudorina elegans*) represent an intermediate stage in this transition. The **Pan/early hominin social protocell** is a pre-ETI starting point exhibiting only pre-adapted evolutionary individuality. **Early Homo** represents the social protocell once it is undergoing an ETI. **Homo** represents the socient, which is the proposed result of the cultural ETI: an integrated cultural unit that fulfills most of the individuality criteria.

For each applied criterion there is also a theoretical expectation that may be argued from the standpoint of the evolutionary trajectory that is invoked. For an egalitarian ETI beginning from "a protocellular situation," we may predict as follows:

At the social protocell stage, **informational uniqueness should be fulfilled** at least to some extent since this is necessary for selectable variation (this is also referred to as "between-group variation"). Social protocells are also expected to **not be informationally homogenous**. Many types of lower-level entities coexist stably within the social protocell as they do not compete directly (e.g., traditions

⁷ We here need to keep track of the difference and linkages between (1) the biological fitness contribution that a traditional strategy may have to its carrier, and (2) the fitness of the tradition itself in terms of its likelihood of spreading between carriers.

Table 2 Comparative summary of the application of criteria for evolutionary individuality (see also corresponding sections)

	Eukaryotic cell	Colonial organism that develops clonally	Multicellular organism	<i>Pan/early</i> hominin—social protocell not undergoing an ETI	Early <i>Homo</i> —social protocell undergoing an ETI	<i>Homo</i> —proposed sociot organism
Spatial/temporal boundaries	Yes. The cell is a cohesive whole and arises in a fission event. It dies in fission or by other causes	Sometimes. If the cells can leave the group and reproduce on their own, then spatial boundaries are absent	Yes. The organism is a cohesive whole, with spatial boundaries, birth, and death	Yes. The primary boundary is in a social space that may or may not correspond to non-overlapping territories. Like cells, communities arise and die in community fission events	Yes, see <i>Pan/early hominin</i>	Yes, see <i>Early Homo</i>. Boundaries gain internal structure, and their cultural control today must reasonably have emerged across the Pleistocene, although details and timing are hard to determine
Informational uniqueness	Yes	Yes	Yes	Sometimes. Boundaries prevent horizontal information spread, thereby preventing uniformity in group composition. Degree of informational uniqueness also depends on the fidelity of social protocell heredity	Yes, see <i>Pan/early hominin</i>. Informational uniqueness is further promoted by the emergence of institutions, which hampers the transfer of culture between units	Yes, see <i>Early Homo</i>. Informational uniqueness may have increased over time as institutions became more complex
Informational homogeneity	No. Egalitarian ETIs do not exhibit informational homogeneity. The genes contained within the eukaryotic cell contain different information	Yes. Since development is clonal, all cells are genetically identical and arise from a unicellular stage. Informational homogeneity is an outcome of a fraternal ETI	Yes. Since development is clonal, all cells are genetically identical and arise from a unicellular stage. Informational homogeneity is an outcome of a fraternal ETI	No. Traditions specialized in different functions are not underpinned by the same information	No. Egalitarian ETIs do not exhibit informational homogeneity. The cultural components of the sociot are underpinned by different information	No. See <i>Early Homo</i>
Indivisibility	Yes. The components of an eukaryotic cell are part of an integrated cellular system	Sometimes. In some species, cells can leave the group and reproduce on their own; for other species, the colony stays together throughout development	Yes. Due to the single-cellular stage and specialized reproductive cells, the cellular level is indivisible. Higher-levels parts (tissue, organ, etc.) are functionally integrated and lack separate fitness	No. The collection of traditions contained within a community may be divided since they are not codependent	Sometimes. The incipient sociot organization of the social protocell is increasingly indivisible . Dependencies between traditions may arise, resulting in the evolution of institutions	Yes. The system of cultural components is a functionally integrated whole. Components are not viable on their own and their contribution to fitness (sociot and hominin) depends on their coadapted cultural context. Fitness has been exported to the new higher level of selection

Table 2 (continued)

	Eukaryotic cell	Colonial organism that develops clonally	Multicellular organism	<i>Pan/early hominin</i> —social protocell not undergoing an ETI	Early <i>Homo</i> —social protocell undergoing an ETI	<i>Homo</i> —proposed sociot organism
Group-level adaptations	Yes. Eukaryotic cells have numerous adaptations, including membrane-bound organelles and meiotic reproduction	Yes. Generally including cell number control, with the specific adaptations dependent upon the species in question	Yes, and increasing relative to undifferentiated ancestors. These include cellular differentiation, tissue specialization, etc., with specific adaptations depending on species	No, or marginally. While under-researched in primates, animal traditions do not appear to possess emergent properties that arise from interactions with other traditions in other contexts	Increasingly from initial low level. The integration and differentiation of institutions, originally around carnivory, is driven by selection on the cultural system as a whole	Yes. Human culture is highly integrated, and institutions appear to be group-level adaptations
Evolutionary division of labor	Yes. An integrated cellular system consists of complementary and specialized intracellular components	No. Cells are undifferentiated and unspecialized	Yes. Cells are differentiated into cell types with different functions and differential expression of a uniform informational underpinning	No. Traditions have different domains but are not integrated, so there is no higher-level labor to divide. Division of labor between components of the traditions themselves is explainable by cognitive processes	Increasingly from initial low level. Integrated and internally organized institutions should emerge at the outset of ETI and this appears to have happened early in the evolution of <i>Homo</i>	Yes. An integrated cultural system consists of complementary and specialized cultural components, e.g., in domains like mobility, tool production, hunting, sharing, comfort, etc
Multilevel selection 2	Yes. Eukaryotic cells have emergent properties, are integrated, and form lineages in which genomes both survive and reproduce as a whole	Sometimes. Depending on the organism's traits (e.g., presence of obligate coloniality, emergent or aggregative survival and reproduction), MLS1 or MLS2 may apply	Yes. Early-onset group selection by close kinship between clonally reproducing protists	No, or very weakly. To the extent that selection acts, it acts on the level of traditions. Integrated systems of traditions are absent (or at least marginal and undetected so far)	Increasingly. Carnivory provides a target around which increasingly sophisticated and integrated cultural systems can form. These cultural systems have emergent properties and can give rise to lineages of systems	Yes. The fate of higher-level entities (sociot as integrated cultural systems) is of primary interest: cultural systems have emergent properties and survive and reproduce as wholes

aimed at different foraging tasks). A social protocell is **divisible** in the sense that the fitnesses of lower-level entities do not depend on the presence of other types of such entities. Although internally heterogenous, the lower-level entities (individual traditions) are not initially organized into systems and exhibit **neither division of labor nor group-level adaptations** at the outset. On the same account, groups of traditions in social protocells **do not exhibit MLS 2**. The social protocell is by definition expected to exhibit **temporal and spatial boundaries**.

Beyond the protocell, an ETI should lead to all these criteria being fulfilled to an increasing degree as a result of selection for evolutionary individuality. **The exception is informational homogeneity**, whose functional effects will be achieved by the evolution of other conflict modifiers. The multicellular organism—our example of a fraternal ETI (Table 2)—should, on the other hand, be expected to also exhibit informational homogeneity (see *Informational homogeneity*).

We find that our assessments match up with theoretical expectations, as well as with the biological examples that we used for comparison. Our findings are consistent with the SPH hypothesis that a cultural evolutionary individual emerged as the outcome of an ETI that may be described as egalitarian.

Discussion

We tested the SPH by subjecting it to a range of criteria developed to identify evolutionary individuals (Clarke 2013; Hanschen et al. 2017). Many of these criteria correspond to mechanisms that promote selection at higher levels (Clarke 2013; Hanschen et al. 2017). We examined *Pan* as a proxy for early hominins, evidence of culture in early *Homo*, and later *Homo* cultural communities. We found that later *Homo* cultural systems satisfied more individuality criteria than did those of early *Homo*, and that early *Homo* cultural systems in turn satisfied more individuality criteria than did *Pan*.

Taken together, our analyses indicate that evolutionary individuality arose, and subsequently increased, in cultural systems during the evolution of *Homo*. Since the features that we have tested for are unlikely to arise and become highly developed for other reasons, this supports the hypothesis that deep-seated similarities exist between the evolutionary provenances of human culture and biological organisms.

But the idea that human culture would have more than superficial similarities with biological organisms is clearly controversial in biology as well as social science (see, e.g., Dunn 2016, pp. 11–31 for a review). We will therefore end by discussing the evolutionary individuality of human culture in the context of four salient differences between human culture and biological organisms.

Let us first briefly remark on what sort of similarities and differences the SPH should lead us to expect. The SPH implies that human cultural systems and biological organisms represent outcomes of the same type of evolutionary process—an ETI—operating in two radically different substrates, namely socially learned behavior and biochemistry (i.e., in the spirit of “general” or “universal” evolutionary theory; see, e.g., Campbell 1974; Hull 1980; Dawkins 1983, 1992; Czikó 1995; Aldrich et al. 2008; Andersson 2008). The expectation is thereby that differences in outcome are attributable to differences in substrate rather than to fundamental differences in the evolutionary process.

First Difference: Are Cultural Communities More Like an Ecosystem?

Recent societies are more often likened with ecosystems than with organisms, and ecosystems would not meet the individuality criteria discussed in the present article (Huneman 2014). Like ecosystems, recent societies exhibit the whole range of ecological relations, including competition, neutralism, parasitism, commensalism, and amensalism (see, e.g., Sandén and Hillman 2011). This raises the question as to whether the SPH overstates the similarity between cultural communities and biological organisms.

The time frame of the sociont (and of this study) is important in this context. The SPH places the base of the ETI at circa 2.0–2.5 Mya, at which time (and earlier) face-to-face coordinated social communities, strongly bounded upward in size by cognitive capacity (e.g. Dunbar 1993; Hill and Dunbar 2003), were the top level of social and cultural organization. This seems to have remained the case until some 50–100 kya when larger and more aggregated social units arose and became dominant during the Late Pleistocene (e.g. Moffett 2013, 2019; see also *Spatial/Temporal boundaries*). During the Holocene, cultural and social organization kept expanding dramatically in level upon level.

The sociont coincides with ancestral cultural communities of the older and smaller style. The more recent and larger aggregates would have required institutions extending between and above the level of the sociont to handle inter-community conflicts (Gat 2010; Wilson 2013). Embedded in such institutions, the original sociont would need to adapt to new and changed roles. These aggregated cultural units would also be much larger, and there would be fewer of them, which would inhibit group selection on levels above that of the sociont (Traulsen and Nowak 2006). If anything, selection on cultural groups would thereby have waned in importance in an increasingly fluid multilevel organization, with less institutional checks on non-cooperative interactions (a “wicked” system; see Andersson and Törnberg 2018; Andersson et al. 2014b).

There is no reason to think that these more recent higher- and multilevel societies would be organized in the same way as the sociot components that they once emerged from. The suitability of ecological models to recent society thereby does not contradict the suitability of an organismal model for ancient societies. The first difference thereby primarily stands out on a comparison between *recent* human societies and biological individuals.

Second Difference: Internally Generated Adaptive Traits

While biological adaptation mainly operates on randomly generated changes in genes and genotypes, adaptive traits in cultural communities (including heritable features) frequently arise by internal innovation. Hominin creativity, and trial-and-error on a fine level of resolution, reacts much more rapidly than selection could have worked on variation on the sociot level. This internal and goal-directed nature of some cultural change raises the question: to what extent do we need cultural group selection to explain cultural adaptation?

Let us look at cultural evolution in action in more detail. High-resolution examples are hard to find in the deep past, so we will consider the emergence of a major institution across the interface between the late Epi-Paleolithic and the earliest Neolithic. Stiner et al. (2014, 2021; Munro et al. 2018) have described in detail how sheep domestication arose over the course of more than a millennium at Aşıklı Höyük (AH; Central Anatolia) —transitioning from management of wild populations, to fully domesticated animals alternately penned and herded to distant pastures (see also Abell et al. 2019). What Stiner and colleagues describe is a multigenerational process where novel solutions, in an iterative and cascading manner, produce new problems to solve and opportunities to pursue. For example, penning sheep within the settlement reduces losses to predators, but creates additional problems, such as with pests. These cascades of change propagate through society as a whole, leading in the end not only to a new institution, but to the integration of this institution as a functional component of the internal organization of the cultural community as a whole.

Several generalizable observations can be made in this example. First, humans here engage collectively in a dynamical and creative innovation process where solving problems and pursuing opportunities generates variation in cultural components, leading to cascades of transformations, and thereby to new problems and opportunities (Andersson et al. 2014a; Lane 2016). Second, while this new institution arises as a result of human problem-solving capabilities, there is no evidence (nor reason to expect) that ideas about the outcome—an integrated system of cultural knowledge and practices making up a pastoral economy—guided the actions taken (which applies to

domestication generally; see, e.g., Zeder 2012, 2015). Humans here built an institution they cannot possibly have understood, they integrated it into a larger cultural system (that they would have understood even less), and it still worked splendidly; see also Lansing's (1987) description of Balinese rice growing communities. Third, and most importantly, the cumulative innovative steps taken *did not* represent the selective replacement of sociot variants in a larger population, nor is it reasonable to believe that populations of variant institutions were maintained simultaneously within the community.

One possible SPH interpretation would be to view this (at least partly) as a developmental rather than evolutionary process. That is, to see innovation as the development of societal organization via a process that *in turn* is based on heritable cultural information, such as via what Heyes (2018) describes as “cognitive gadgets”; see also Ardila (2018), with writing and mathematics used as examples. Compared with biological counterparts, the degree of sociot developmental plasticity would truly be exceptional, but, then again, the affordances of a cultural organism would be radically different from those of a biological organism. Mechanisms for altering phenotypic expression via a flexible developmental process have clearly been strongly selected for, and played an important role, in biological evolution (e.g., West-Eberhard 2003; Sterelny 2011; LaFreniere and MacDonald 2013). In the sociot context, Andersson and Törnberg (2019) argued that one of the major advantages of an environmentally responsive and integrated cultural system could have been to leverage the high flexibility that generally is a trademark of great ape behavior (e.g., Ungar et al. 2006; Malone et al. 2012); see also, e.g., Fogarty et al. (2015) and Fuentes (2017b). That proposition dovetails with the “variability selection hypothesis” (Potts 1998, 2012; Grove 2011b, 2011a; Maslin et al. 2014, 2015) which argues that high levels of environmental variation during the early Pleistocene would have strongly favored any ability to rapidly reconfigure one's behavior.

If so, the main targets of sociot selection and adaptation would not have been the detailed manifestations of culture, but something more toward the image of cognitive gadgets (Ardila 2018; Heyes 2018); i.e., fundamental systems that underpin our capacity to adapt culturally by governing how we think and by harnessing the capacity of our large brains (Andersson and Törnberg 2019, p. 86). For example, the people at Aşıklı Höyük would have been in a position to embark on their transformation into a Neolithic society (once facing Holocene conditions; e.g., Richerson et al. 2001) because earlier cultural communities (and hominins) in that lineage had accumulated, across hundreds and thousands of millennia, a richer and richer system of basic cognitive tools for dealing successfully with environmental change in general.

Basic and stably present systems like these would over time be expected to coevolve with *Homo* to form “hybrid” systems, whose genetic and cultural elements would become so closely intertwined that clearly classifying them as either genetic or cultural is impossible. Language could provide the most salient example of such a system. Language is underpinned by genetic adaptations (e.g., Mozzi et al. 2016), but we learn languages, and they evolve also in their own right (e.g., Croft and Cruse 1987; Mufwene 2001; Greenhill et al. 2010). Language is also far more than just a means of communication. It structures the complexities of human cognition and psychology, which, essentially, makes it an operating system for our large and expensive brains. It is widely agreed that language must have evolved gradually from a protolanguage. But while a protolanguage would have been useful, or even necessary, for early *Homo* (e.g., Bickerton 2009; Bickerton and Szathmáry 2011; in the context outlined in Fig. 2), it is still hard to account for its original emergence in terms of benefits accruing to individual hominins. The sociot would be expected to drive the evolution of precisely this type of complex and shared cultural systems that would become part of the selective environment of *Homo*.

Third Difference: Lower-Level Selection

Cultural evolution as selection acting on populations of variants of cultural components that arise and spread within cultural communities is a well-researched and central theme in cultural evolutionary studies (Cavalli-Sforza and Feldman 1981; Boyd and Richerson 1985, 2005; Durham 1992; Mesoudi 2007, 2011). This raises the question of how much selection remains in operation also on lower-level cultural components, and how this squares with the notion that lower-level selection would have been suppressed during an ETI?

We have identified multiple mechanisms that inhibit lower-level selection, as many of the individuality criteria correspond to mechanisms that promote higher-level selection and/or reduce lower-level selection (Clarke et al. 2013; Hanschen et al. 2017). But egalitarian ETIs never eliminate lower-level selection completely (see also the “Analysis” of the “Multilevel Selection 2” section). Lower-level selection could still be occurring within the sociot, as it does within biological individuals that have emerged in egalitarian ETI. For example, selection on genes is not completely suppressed in cells (see, e.g., Ågren 2014).

Lower-level selection may also belong to sociot mechanisms adapted specifically to increase the capacity to respond to the environment (i.e., as part of the types of mechanisms discussed as developmental in the previous section); see also Ziman (2000). Employing selection in such a role would not lack precedents. There are two major examples of biological organs that operate as Darwinian systems based on staged

and adapted implementations of “blind-variation-selective-retention” (BVSR; Campbell 1960): the adaptive immune system and the brain (e.g., Jerne 1955; Changeux 1985; Michod 1988; Edelman 1993; Fernando and Szathmáry 2010; Müller et al. 2018). The function of these organs is precisely to provide the biological organism with capabilities for responding to the environment on timescales that are too short for genetic adaptation; for example, creativity, learning, and the ability to survive the onslaught of pathogenic microorganisms with much shorter generation times.

Fourth Difference: Boundaries and Manifestation

Biological individuals tend to be physically cohesive, and individuality criteria such as *spatial/temporal boundaries* and *indivisibility* are easy to interpret in terms of physical boundaries. The sociot, however, must be imagined largely in other spaces, such as social and ideational spaces. Are such boundaries equivalent with those in biology? We described sociot boundaries in the “Spatial/Temporal Boundaries” section but expand our discussion here with a tentative description of how a sociot would manifest itself—along with *Homo*—in cultural communities (see also the “Second Difference: Internally Generated Adaptive Traits” section).

On the basis of the analysis in this article, we propose that the phenotypic manifestation of the sociot may be pictured as a stationary and organized pattern of behaviors, cultural products, and environmental modifications—coincident with, and maintained by, but not identical to, a social community of hominins. It would be generated and maintained by the dynamical and parallel expression (by hominins) of cultural components, most of which are likely tacit. This emergent pattern would unfold in time and space as the expression of cultural components regulated the expression of other components (within and between brains) via social interactions, cultural products, and environmental modifications. Expressed cultural components would act by modulating hominin behavior via psychology, cognition, and metacognition.

The stationary structure of this dynamical pattern may be conceptualized schematically (in the manner of an organizational chart) as a nested hierarchy of functional subsystems – such as for hunting, fishing, tracking, knapping, pyrotechnology, but also strategies for things like teaching, distributing resources, resolving conflicts, and so on. This organization may be unpacked all the way into the individual brain, where culture interfaces with our psychology and cognition. In terms of extent, this system reaches only as far as the social interactions of its carriers—i.e., it has a boundary, and, since culture shapes social interactions, the nature and extent of this boundary is evolvable as a part of the system itself.

Moreover, the above description of the sociot potentially dovetails with other models of the dynamics and organization of culture within as well as between cognitive agents.

For example, Heyes (2018) describes cognitive gadgets as not only functional but also regulatory systems, acting within the brain to form adapted systems from highly domain-general innate components. The autocatalytic network model by Gabora and Steel (Gabora and Steel 2017, 2020) sees learning and creativity as a result of self-organization in mental representation networks. They suggest an extension of these dynamics to the social level, which could pertain to the above envisioned intra-sociot dynamics, whose mechanisms would then be shaped by sociot evolution (or mutualistic sociot-hominin coevolution.)

Models depicting emergent "group cognition" in networked human cognitive nodes, organized and mediated by culture, have been proposed by several authors (e.g., Grove and Coward 2008; Coward and Grove 2011; Gallagher 2013; Muthukrishna and Henrich 2016; Read 2020), including models where culture and its products themselves are depicted as part of an "external mind" (Clark and Chalmers 1998; Menary 2010); see Theiner (2014) for a review. Cultural niche construction focuses on complex causal feedback loops between cultural behavior and persistent environmental features (Laland and Brown 2006; Smith 2007; Laland and O'Brien 2015), and networked, recombining and cascading features in general are central in many theories of innovation in modern and ancient sociotechnical systems (see, e.g., Hughes 1986; Geels 2002; Schiffer 2005; Wimsatt and Griesemer 2007; Andersson et al. 2014a; Lane 2016; d'Errico and Colagè 2018).

The description also recalls several models of biological innovation and organization in a recent family of models often referred to as the "extended evolutionary synthesis" (e.g., Pigliucci and Müller 2010; Feldman et al. 2015; Jaeger et al. 2015); for its applications to culture see, for example, Andersson et al. (2014a), Fuentes (2016, 2017a), Smith et al. (2018), and Zeder (2018). Gene regulatory networks in biological development (e.g., Arthur 2011) exhibit dynamical and evolutionary similarities with socio-technical innovation (e.g., Erwin and Krakauer 2004). Also, the extension of genes, via social interactions, to group-level adaptations in social insects (via tactile and chemical signals as well as by sensing of persistent modifications of the environment) leads to the formation of a biological organismal unit that also challenges the view of organisms as physically bounded and contiguous entities (Dorigo et al. 2000; Queller and Strassmann 2009; Kennedy et al. 2017).

Future Directions

Many important issues could not be covered in this article. For example, while evolutionary individuality suggests that competition *can* take place on the group level, we have not discussed the issue of how competition actually would happen. Discussions of group-level competition provided by Smaldino (2014) and (Richerson et al. 2016) apply largely also to the SPH (even if the delimitation between genetic and cultural information is unclear in those treatments), but the issue must be expanded on in future work.

Formal modelling may help to clarify the role of group-level competition, and, in general, to investigate what assumptions are needed for the postulated links between the entities (e.g., in terms of fitnesses of traditions, hominins, and sociots) to operate as claimed, and to explore phenomena in populations and over time.

Future work should also examine derived human traits such as cooperation, altruism, and language in the context of the sociot and the proposed mutualistic partnership with *Homo*. This relation must be worked out theoretically, via models, via empirical examples, as well as via revisiting frameworks such as those mentioned in the previous section, including not least models of coevolution between *Homo* and cultural systems (Durham 1992; Herrmann et al. 2007; Smith 2007; Laland and O'Brien 2015; Fernandes and Woodley of Menie 2017; Hare 2017; Colagè and d'Errico 2018).

It would have high diagnostic value if features of culture and/or humans could be demonstrated to be adapted to evolutionary meta-functions of the sociot (see above). Generally, the prediction that culture was shaped by the same processes that generate organismic form in biology should be further explored. The concept of "organismality" (e.g., Queller and Strassmann 2009; West and Kiers 2009; Strassmann and Queller 2010) here represents an attempt to understand what it means to be an organism in the abstract, and it may serve as a useful starting point.

As argued in "The Social Protocell" section and the "Analysis" in the "Informational Uniqueness" section, we may imagine high-fidelity inheritance of "proto-institutions" as systems of functions *before* high-fidelity social learning enabled cumulative evolution of the processes underpinning the functions. This requires only the functions to be stable, which may be the case also if they are underpinned by processes that are constantly reinvented rather than inherited (i.e., emulated rather than imitated). Such a proto-cultural phase would require laxer assumptions about early hominin cognitive evolution than a direct move from emulation to an inheritance system based on high-fidelity imitation (see, e.g., Shea 2009).

Finally, do marginal proto-institutions exist in *Pan*? If they do, they could provide simple analogs for a proto-cultural stage of the evolution of human culture. Such systems may have emerged and remained marginal if combinations of stable socially learned behaviors produced adaptive effects on the community level, but without the open-endedness argued to apply to big game carnivory. Read (2012, pp. 99–104), for example, describes substantial intraspecies regional variability in collective group behavior in *Pan*, such as in how border patrols are organized, without evident genetic differences to explain this variation.

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Glossary

Tradition Patterns of thinking and behavior (e.g., food preferences, foraging strategies, tools, social customs, etc.) that persist across generations due to social learning. Widespread among animals, traditions are foremostly adapted by creative problem solving, but with a potential for limited Darwinian selection arising from repeated cycles of learning and application of variant instances held by different learners.

Cultural component Human cultural systems cannot generally be resolved into identifiable discrete traditions. To denote the parts of cultural systems more generally we speak instead of cultural components. For particularly large and integrated cultural components we speak of institutions.

Culture We reserve the term culture for integrated and adapted systems of cultural components (originally traditions.) Only the genus *Homo* supports culture in this view, and “an animal culture” is simply a collection of stand-alone traditions without significant interactions between them (see also Galef 1992; Read and Andersson 2019).

Social protocell A system of independent features of social group behavior and learning that together creates (as a side effect) a potential for sets of animal traditions to be selected together. Like the biological protocell – which plays a key role in the origin of cellular life – it imparts essential meta-evolutionary functions (reproduction, heredity, and containment) to a potential future group-level evolutionary individual.

Evolutionary individual An evolutionary individual is an integrated higher-level unit of selection and adaptation in which selection of lower levels components is restricted. For instance, the multicellular organism is an individual as selection at the lower level of the cell is restricted.

Sociont The SPH proposes that groups of tradition evolved into a new kind of group-level cultural entity, termed a sociont, comprised of integrated institutional components (i.e., institutions.) The potential that evolving institutions may improve the efficiency of meta-evolutionary functions is the evolution of evolutionary individuality that we investigate in this paper.

Cultural community A social community of animals (in practice *Homo*) whose behaviors are largely governed by a cultural system that they enact, maintain, and transmit. Following the proposition by Andersson and Törnberg (2019), it is seen here as an obligate mutualistic partnership between *Homo* and sociont.

Evolutionary Transition in Individuality (ETI) Groups of individuals become a new kind of individual. In an ETI, formerly independent lower-level evolutionary individuals are integrated into a new group-level evolutionary individual. The lower-level entities cease, wholly or partly, to be evolutionary individuals in their own right – such as cells in a multicellular organism or organelles (e.g., mitochondria) in eukaryotic cells.

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References

- Abell JT, Quade J, Duru G, Mentzer SM, Stiner MC, Uzundurum M, Özbaşaran M (2019) Urine salts elucidate early Neolithic animal management at Aşikli Höyük, Turkey. *Sci Adv* 5(4).
- Ågren JA (2014) Evolutionary transitions in individuality: insights from transposable elements. *Trends Ecol Evol* 29(2):90–96
- Aldrich HE, Hodgson GM, Hull DL, Knudsen T, Moky J, Vanberg VJ (2008) In defence of generalized darwinism. *J Evol Econ* 18:577–596
- Allen JA (2019) Community through culture: from insects to whales: how social learning and culture manifest across diverse animal communities. *BioEssays* 41(11):1–8
- Andersson C (2008) Sophisticated selectionism as a general theory of knowledge. *Biol Philos* 23(2):229–242
- Andersson C (2011) Palaeolithic punctuations and equilibria: did retention rather than invention limit technological evolution? *Palaeo-Anthropology* 243–259
- Andersson C (2013) Fidelity and the emergence of stable and cumulative sociotechnical systems. *PalaeoAnthropology* 2013:88–103
- Andersson C, Törnberg A, Törnberg P (2014a) An evolutionary developmental approach to cultural evolution. *Curr Anthropol* 55(2):154–174
- Andersson C, Törnberg A, Törnberg P (2014b) Societal systems: complex or worse? *Futures* 63:145–157
- Andersson C, Törnberg P (2018) Wickedness and the anatomy of complexity. *Futures* 95:118–138
- Andersson C, Törnberg P (2019) Toward a macroevolutionary theory of human evolution: the social protocell. *Biol Theory* 14(2):86–102
- Ardila A (2018) Historical development of human cognition. Springer Nature, Singapore
- Arthur W (2011) Evolution: a developmental approach. Wiley-Blackwell, Chichester
- Bar-Yosef O, Wang Y (2012) Paleolithic archaeology in China. *Annu Rev Anthropol* 41:319–335

- Beinhocker E (2006) The origin of wealth: evolution, complexity, and the radical remaking of economics. Harvard Business School Press, Boston
- Bickerton D (2009) Adam's tongue. Macmillan Press, New York
- Bickerton D, Szathmáry E (2011) Confrontational scavenging as a possible source for language and cooperation. *BMC Evol Biol* 11(1):261
- Blasco R, Rosell J, Domínguez-Rodrigo M, Lozano S, Pastó I, Riba D et al (2013) Learning by heart: cultural patterns in the faunal processing sequence during the middle pleistocene. *PLoS One* 8(2). <https://doi.org/10.1371/journal.pone.0055863>
- Blegen N (2017) The earliest long-distance obsidian transport: evidence from the ~200 Ka Middle Stone Age Sibilo School Road Site, Baringo, Kenya. *J Hum Evol* 103:1–19
- Boesch C (2012) Wild cultures: a comparison between chimpanzee and human cultures. Cambridge University Press, New York
- Boesch C, Crockford C, Herbinger I, Wittig R, Moebius Y, Normand E (2008) Intergroup conflicts among chimpanzees in tai national park: lethal violence and the female perspective. *Am J Primatol* 70(6):519–532
- Boesch C, Tomasello M (1998) Chimpanzee and human cultures. *Curr Anthropol* 39(5):591–614
- Boyd RT, Richerson PJ (1985) Culture and the evolutionary process. University of Chicago Press, Chicago
- Boyd RT, Richerson PJ (1987) The evolution of ethnic markers. *Cult Anthropol* 2(1):65–79
- Boyd RT, Richerson PJ (1992) Punishment allows the evolution of cooperation (or anything else) in sizable groups. *Ethol Sociobiol* 13(3):171–195
- Boyd RT, Richerson PJ (2000) Memes: universal acid or a better mouse trap? In: Aunger R (ed) *Darwinizing culture: the status of memetics as a science*. Oxford University Press, Oxford, pp 143–162
- Boyd RT, Richerson PJ (2005) Not by genes alone. University of Chicago Press, Chicago
- Boyd RT, Richerson PJ (2010) Transmission coupling mechanisms: cultural group selection. *Philosophic Transact Royal Soc B: Biol Sci* 365(1559):3787–3795
- Brooks AS, Yellen JE, Potts R, Behrensmeier AK, Deino AL, Leslie DE et al (2018) Long-distance stone transport and pigment use in the earliest Middle Stone age. *Science* 360(6384):90–94
- Buskell A, Enquist M, Jansson F (2019) A systems approach to cultural evolution. *Palgrave Commun* 5(1):1–15
- Buss LW (1987) The evolution of individuality. Princeton University Press, Princeton, NJ
- Campbell DT (1960) Blind variation and selective retention in creative thought as in other knowledge processes. *Psychol Rev* 67(6):380–400
- Campbell DT (1974) Evolutionary epistemology. In: Schilpp PA (ed) *The philosophy of Karl Popper*, Open Court, pp 413–463
- Cavalli-Sforza LL, Feldman MW (1981) Cultural transmission and evolution. Stanford University Press, Stanford
- Changeux J-P (1985) Neuronal man: the biology of mind. Oxford University Press, New York
- Chudek M, Henrich J (2011) Culture-gene coevolution, norm-psychology and the emergence of human prosociality. *Trends Cogn Sci* 15(5):218–226
- Clark A, Chalmers D (1998) The extended mind. *Analysis* 58(1):7–19
- Clarke E (2013) The multiple realizability of biological individuals. *J Philosophy* 110(8):413–435
- Clarke E (2014) Origins of evolutionary transitions. *J Biosci* 39(2):303–317
- Clay Z, Tennie C (2018) Is overimitation a uniquely human phenomenon? Insights from human children as compared to bonobos. *Child Dev* 89(5):1535–1544
- Colagè I, d'Errico F (2018) Culture: the driving force of human cognition. *Top Cogn Sci* 12:654–672
- Coward F, Grove M (2011) Beyond the tools: social innovation and hominin evolution. *PaleoAnthropology* 111–229
- Croft W, Cruse DA (1987) Cognitive linguistics. Cambridge University Press, Cambridge
- Csibra G, Gergely G (2009) Natural pedagogy. *Trends Cogn Sci* 13(4):148–153
- Csibra G, Gergely G (2011) Natural pedagogy as evolutionary adaptation. *Philosophic Trans Royal Soc London. Series B Biol Sci* 366(1567):1149–57
- Cziko G (1995) Without miracles. MIT Press, Cambridge
- d'Errico F, Colagè I (2018) Cultural exaptation and cultural neural reuse: a mechanism for the emergence of modern culture and behavior. *Biol Theory* 13(4):213–227
- Damuth J, Lorraine Heisler I (1988) Alternative formulations of multilevel selection. *Biol Philos* 3(4):407–430
- Dawkins R (1983) Universal Darwinism. In: Bendall DS (ed) *Evolution from molecules to man*. Cambridge University Press, Cambridge, pp 403–425
- Dawkins R (1992) Universal biology. *Nature* 360(6399):25–26
- de Waal FBM (2013) Animal conformists. *Science* 340:437–438
- Dorigo M, Bonabeau E, Theraulaz G (2000) Ant algorithms and stigmergy. *Futur Gener Comput Syst* 16(8):851–871
- Dugatkin LA (2002) Cooperation in animals: an evolutionary overview. *Biol Philos* 17(4):459–476
- Dunbar RIM (1992) Neocortex size as a constraint on group size in primates. *J Hum Evol* 22(6):469–493
- Dunbar RIM (1993) Coevolution of neocortical size, group size and language in humans. *Behavioral and Brain Sciences* 16:681–735
- Dunbar RIM (1998) The social brain hypothesis. *Evol Anthropol* 6(5):178–190
- Dunn MB (2016) Reviving the organismic analogy in sociology: human society as an organism. Dissertation, University of California Riverside
- Dunstone J, Caldwell CA (2018) Cumulative culture and explicit meta-cognition: a review of theories, evidence and key predictions. *Palgrave Commun* 4(1):1–11
- Durand PM, Michod RE (2010) Genomics in the light of evolutionary transitions. *Evolution* 64(6):1533–1540
- Durham WH (1992) Coevolution - genes, culture, and human diversity. Stanford University Press, Stanford
- Edelman GM (1993) Neural Darwinism: selection and reentrant signaling in higher brain function. *Neuron* 10(2):115–125
- Eigen M, Schuster P (1977) The hypercycle. a principle of natural self-organization. *Naturwissenschaften* 64:541–565
- Erwin DH, Krakauer DC (2004) Insights into innovation. *Science* 304:1117–1119
- Féblot-Augustins J (1997) La Circulation Des Matières Premières Au Paléolithique: Synthèse Des Données: Perspectives Comportementales. In: *Études et recherches archéologiques de l'Université de Liège*, 75. Service de Préhistoire, Université de Liège, Liège
- Feldblum J, Manfredi S, Gilby I, Pusey AE (2018) The timing and causes of a unique chimpanzee community fission preceding Gombe's Four Years' War. *Am J Phys Anthropol* 166:730–744
- Feldman MW, Laland KN, Uller T, Sterelny K, Müller GB, Moczek A, Jablonka E, Odling-Smee J (2015) The extended evolutionary synthesis: its structure, assumptions and predictions. *Proc Royal Society B Biol Sci*. <https://doi.org/10.1098/rspb.2015.1019>
- Fernandes HBF, Michael A, Woodley of Menie (2017) Darwin's Unfinished Symphony: How Culture Made the Human Mind. *Anim Behav* 133:207–208
- Fernando C, Szathmáry E (2010) Natural selection in the brain. In: *Towards a theory of thinking - building blocks for a conceptual framework*. Springer, Berlin/Heidelberg, pp 291–332

- Fogarty L, Creanza N, Feldman MW (2015) Cultural evolutionary perspectives on creativity and human innovation. *Trends Ecol Evol* 30(12):736–754
- Foley RA, Mirazón Lahr M (2011) The evolution of the diversity of cultures. *Philosophic Trans Royal Soc London. Series B, Biol Sci* 366(1567):1080–89
- Fuentes A (2016) The extended evolutionary synthesis, ethnography, and the human niche: toward an integrated anthropology. *Curr Anthropol* 57(S13):S13–26
- Fuentes A (2017a) Human niche, human behaviour, human nature. *Interface Focus* 7(5):1–42
- Fuentes A (2017b) The creative spark: how imagination made humans exceptional. Penguin, New York
- Furuichi T (1987) Sexual swelling, receptivity, and grouping of wild pygmy chimpanzee females at Wamba, Zaïre. *Primates* 28(3):309–318
- Gabora L, Steel M (2017) Autocatalytic networks in cognition and the origin of culture. *J Theor Biol* 431:87–95
- Gabora L, Steel M (2020) Modeling a cognitive transition at the origin of cultural evolution using autocatalytic networks. *Cogn Sci* 44(9):1–29
- Galef BG (1992) The question of animal culture. *Hum Nat* 3(2):157–178
- Galef BG, Laland KN (2005) Social learning in animals : empirical studies and theoretical models. *Bioscience* 55(6):489–499
- Gallagher S (2013) The socially extended mind. *Cogn Syst Res* 25–26:4–12
- Gánti T (1975) Organization of chemical reactions into dividing and metabolizing units: the chemotons. *BioSystems* 7(1):15–21
- Gánti T (1997) Biogenesis itself. *J Theor Biol* 187(4):583–593
- Gao X (2013) Paleolithic cultures in China: uniqueness and divergence. *Curr Anthropol* 54(SUPPL8)
- Gärdenfors P, Högborg A (2017) The archaeology of teaching and the evolution of *Homo Docens*. *Curr Anthropol* 58(2):188–208
- Gat A (2010) Why war? Motivations for fighting in the human state of nature. In: Kappeler PM, Silk JB (eds) *Mind the gap*. Springer Verlag, Berlin/Heidelberg, pp 197–220
- Geels FW (2002) Technological transitions as evolutionary reconfiguration processes: a multi-level perspective and a case-study. *Res Policy* 31(8–9):1257–1274
- Gergely G, Csibra G (2006) Sylvia's recipe: the role of imitation and pedagogy in the transmission of human culture. In: Enfield NJ, Levinson SC (eds) *Roots of human sociality: culture, cognition, and human interaction*. Berg Press, Oxford, pp 229–555
- Gintis H, van Schaik CP, Boehm C (2015) Zoon Politikon. *Curr Anthropol* 56(3):327–353
- Goodall J (1986) *The chimpanzees of Gombe: patterns of behavior*. Belknap Press, Cambridge, MA
- Greenhill SJ, Atkinson QD, Meade A, Gray RD (2010) The shape and tempo of language evolution. *Proc Royal Soc B Biol Sci* 277(1693):2443–2450
- Grove M (2011a) Change and variability in plio-pleistocene climates: modelling the hominin response. *J Archaeol Sci* 38(11):3038–3047
- Grove M (2011b) Speciation, diversity, and mode 1 technologies: the impact of variability selection. *J Hum Evol* 61(3):306–319
- Grove M, Coward F (2008) From individual neurons to social brains. *Camb Archaeol J* 18(3):387–400
- Grove M, Pearce E, Dunbar RIM (2012) Fission-fusion and the evolution of hominin social systems. *J Hum Evol* 62(2):191–200
- Haidle MN, Bolus M, Collard M, Conard NJ, Garofoli D, Lombard M et al (2015) The nature of culture: an eight-grade model for the evolution and expansion of cultural capacities in hominins and other animals. *J Anthropol Sci* 93:43–70
- Hamilton WD (1964a) The genetical evolution of social behaviour. I. *J Theor Biol* 7:1–16
- Hamilton WD (1964b) The genetical evolution of social behaviour. II. *J Theor Biol* 7:17–52
- Han S, Ma Y (2015) A culture-behavior-brain loop model of human development. *Trends Cogn Sci* 19(11):666–676
- Hanschen ER, Davison DR, Grochau-Wright ZI, Michod RI (2017) Evolution of individuality: a case study in the volvocine green algae. *Philosophy, Theor Pract Biol* 9(20170801)
- Hanschen ER, Davison DR, Zachariah I, Michod RE (2018) Individuality and the major evolutionary transitions. In: *Landscapes of collectivity in the life sciences*. MIT Press, Cambridge, pp 255–268
- Hanschen ER, Shelton DE, Michod RE (2015) Evolutionary transitions in individuality and recent models of multicellularity. In: Ruiz-Trillo I, Nedelcu AM (eds) *Evolutionary transitions to multicellular life*. Springer Science and Business Media, Dordrecht, pp 165–187
- Hare B (2017) Survival of the friendliest: *Homo sapiens* evolved via selection for prosociality. *Ann Rev Psychol* 68(1):annurev-psych-010416–044201
- Harmand S, Lewis JE, Feibel CS, Lepre CJ, Prat S, Lenoble A et al (2015) 3.3-million-year-old stone tools from Lomekwi 3, West Turkana, Kenya. *Nature* 521:310–315
- Haun DBM, Rekers Y, Tomasello M (2012) Majority-biased transmission in chimpanzees and human children, but not orangutans. *Curr Biol* 22(8):727–731
- Haun DBM, Rekers Y, Tomasello M (2014) Children conform to the behavior of peers; other great apes stick with what they know. *Psychol Sci* 25(12):2160–2167
- Henrich J (2004) Cultural group selection, coevolutionary processes and large-scale cooperation. *J Econ Behav Organ* 53:143–162
- Henrich J, Boyd R (2008) Division of labor, economic specialization, and the evolution of social stratification. *Curr Anthropol* 49(4):715–724
- Henrich J, Gil-White FJ (2001) The evolution of prestige: freely conferred deference as a mechanism for enhancing the benefits of cultural transmission. *Evol Hum Behav* 22(3):165–196
- Henrich J, McElreath R (2003) The evolution of cultural evolution. *Evol Anthropol: Issues News Rev* 12(3):123–135
- Herrmann E, Call J, Hernández-Lloreda V, Hare B, Tomasello M (2007) Humans have evolved specialized skills of social cognition: the cultural intelligence hypothesis. *Science* 317(5843):1360–1366
- Heyes CM (2018) *Cognitive gadgets*. Belknap Press, Harvard University, Cambridge
- Hill RA, Dunbar RIM (2003) Social network size in humans. *Hum Nat* 14(1):53–72
- Horner V, Proctor D, Bonnie KE, Whiten A, de Waal FBM (2010) Prestige affects cultural learning in chimpanzees. *PLoS One* 5(5):1–5
- Hughes TP (1986) The seamless web: technology, science, etcetera, etcetera. *Soc Stud Sci* 16:281–292
- Hull DL (1980) Individuality and selection. *Annu Rev Ecol Syst* 11:311–332
- Huneman P (2014) Individuality as a theoretical scheme. II. About the weak individuality of organisms and ecosystems. *Biol Theory* 9(4):374–381
- Jablonka E, Szathmáry E (1995) The evolution of information storage and heredity. *Trends Ecol Evol* 10(5):206–211
- Jaeger J, Laubichler MD, Callebaut W (2015) The comet cometh: evolving developmental systems. *Biol Theory* 10(1):36–49
- Jerne N (1955) The natural-selection theory of antibody formation. *Proc Natl Acad Sci USA* 41(11):67–68
- Jiménez ÁV, Mesoudi A (2019) Prestige-biased social learning: current evidence and outstanding questions. *Palgrave Commun* 5(1):1–12

- Kalan AK, Kulik L, Arandjelovic M, Boesch C, Haas F, Dieguez P et al (2020) Environmental variability supports chimpanzee behavioural diversity. *Nat Commun*. <https://doi.org/10.1038/s41467-020-18176-3>
- Kaufhold SP, Edwin JC, Van L (2019) Why intergroup variation matters for understanding behaviour. *Biol Lett* 15(11)
- Kendal RL, Hopper LM, Whiten A, Brosnan SF, Lambeth SP, Schapiro SJ, Hoppitt W (2015) Chimpanzees copy dominant and knowledgeable individuals: implications for cultural diversity. *Evol Hum Behav* 36(1):65–72
- Kennedy P, Baron G, Qiu B, Freitak D, Helanterä H, Hunt ER et al (2017) Deconstructing superorganisms and societies to address big questions in biology. *Trends Ecol Evol* 32(11):861–872
- Kline MA (2015) How to learn about teaching: an evolutionary framework for the study of teaching behavior in humans and other animals. *Behav Brain Sci* 38:1–70
- Koops K, Schöning C, Isaji M, Hashimoto C (2015) Cultural differences in ant-dipping tool length between neighbouring chimpanzee communities at Kalinzu, Uganda. *Sci Rep* 5(1):12456
- Kuhn SL (2020) The evolution of Paleolithic technologies. Routledge (Taylor Francis), Milton Park
- Kuhn SL, Stiner MC (2006) What's a mother to do? The division of labor among neandertals and modern humans in Eurasia. *Curr Anthropol* 47(6):953–980
- LaFreniere P, MacDonald K (2013) A post-genomic view of behavioral development and adaptation to the environment. *Dev Rev* 33(2):89–109
- Laland KN (2017) The foundations of human cooperation in teaching and imitation. *Spanish J Psychol* 19:E100
- Laland KN, Brown GR (2006) Niche construction, human behavior, and the adaptive-lag hypothesis. *Evol Anthropol* 15:95–104
- Laland KN, O'Brien MJ (2015) Niche construction: implications for human sciences. In: Emerging trends in the social and behavioral sciences: an interdisciplinary, searchable, and linkable resource. Wiley, Hoboken, pp 1–10
- Lane DA (2016) Innovation cascades: artefacts, organization and attributions. *Philosophic Trans Royal Soc London. Series B, Biol Sci* 371(1690):20150194
- Lansing JS (1987) Balinese 'water temples' and the management of irrigation. *Am Anthropol* 89(2):326–341
- Layton R, O'Hara S, Bilsborough A (2012) Antiquity and social functions of multilevel social organization among human hunter-gatherers. *Int J Primatol* 33(5):1215–1245
- Leeuwen V, Edwin JC, Cronin KA, Schütte S, Call J, Haun DBM (2013) Chimpanzees (*Pan troglodytes*) flexibly adjust their behaviour in order to maximize payoffs, not to conform to majorities. *PLoS One* 8(11):1–10
- Leigh EG (2010) The evolution of mutualism. *J Evol Biol* 23(12):2507–2528
- Lewontin RC (1970) The units of selection. *Annu Rev Ecol Syst* 1:1–18
- Lewontin RC (1972) The apportionment of human diversity. *Evol Biol* 6:381–398
- Luncz LV, Boesch C (2014) Tradition over trend: neighboring chimpanzee communities maintain differences in cultural behavior despite frequent immigration of adult females. *Am J Primatol* 76(7):649–657
- Luncz LV, Boesch C (2015) The extent of cultural variation between adjacent chimpanzee (*Pan troglodytes verus*) communities - a microecological approach. *Am J Phys Anthropol* 156(1):67–75
- Lycett SJ, Collard M, McGrew WC (2009) Cladistic analyses of behavioural variation in wild *Pan troglodytes*: exploring the chimpanzee culture hypothesis. *J Hum Evol* 57(4):337–349
- Malone N, Fuentes A, White FJ (2012) Variation in the social systems of extant hominoids: comparative insight into the social behavior of early hominins. *Int J Primatol* 33(6):1251–1277
- Marengo L, Dosi G (2005) Division of labor, organizational coordination and market mechanisms in collective problem-solving. *J Econ Behav Organ* 58(2):303–326
- Maslin MA, Brierley CM, Milner AM, Shultz S, Trauth MH, Wilson KE (2014) East African climate pulses and early human evolution. *Quatern Sci Rev* 101:1–17
- Maslin MA, Shultz S, Trauth MH (2015) A synthesis of the theories and concepts of early human evolution. *Philosophic Trans Royal Soc London. Series B, Biol Sci* 370:1–12
- Maynard-Smith J, Szathmáry E (1993) The origin of chromosomes. I. selection for linkage. *J Theor Biol* 164:437–446
- Maynard-Smith J, Szathmáry E (1995) Major transitions in evolution. W.H. Freeman Press, New York
- McGrew WC (2010) In search of the last common ancestor: new findings on wild chimpanzees. *Philosophic Trans Royal Society B: Biol Sci* 365(1556):3267–3276
- Menary R (2010) The extended mind. In: Menary R (ed) MIT Press, Cambridge
- Mercader J, Barton H, Gillespie J, Harris J, Kuhn SL, Tyler R, Boesch C (2007) 4,300-year-old chimpanzee sites and the origins of percussive stone technology. *Proc Natl Acad Sci USA* 104(9):3043–3048
- Mercader J, Panger M, Boesch C (2002) Excavation of a chimpanzee stone tool site in the African rainforest. *Science* 296:1452–1455
- Mesoudi A (2007) Biological and cultural evolution: similar but different. *Biol Theory* 2(2):119–123
- Mesoudi A (2011) Cultural evolution. University of Chicago Press, Chicago
- Michod RE (1983) Population biology of the first replicators: on the origin of the genotype, phenotype and organism. *Am Zool* 23:5–14
- Michod RE (1988) Book review: Darwinian selection in the brain. *Evolution* 43(3):694–696
- Michod RE (1999) Darwinian dynamics: evolutionary transitions in fitness and individuality. Princeton University Press, Princeton
- Michod RE (2007) Evolution of individuality during the transition from unicellular to multicellular life. *Proc Natl Acad Sci USA* 104 Suppl(suppl_1):8613–8618
- Michod RE, Herron MD (2006) Cooperation and conflict during evolutionary transitions in individuality. *J Evol Biol* 19(5):1406–1409
- Michod RE, Nedelcu AM (2003) On the reorganization of fitness during evolutionary transitions in individuality. *Integr Comp Biol* 43(1):64–73
- Michod RE, Roze D (2001) Cooperation and conflict in the evolution of multicellularity. *Heredity* 86:1–7
- Moffett MW (2013) Human identity and the evolution of societies. *Hum Nat* 24(3):219–267
- Moffett MW (2019) The human swarm. Basic Books, New York
- Mozzi A, Forni D, Clerici M, Pozzoli U, Mascheretti S, Guerini FR et al (2016) The evolutionary history of genes involved in spoken and written language: beyond FOXP2. *Sci Rep* 6:1–12
- Mufwene SS (2001) The ecology of language evolution. Cambridge University Press, Cambridge
- Müller V, de Boer RJ, Bonhoeffer S, Szathmáry E (2018) An evolutionary perspective on the systems of adaptive immunity. *Biol Rev* 93(1):505–528
- Munro ND, Bar-Oz G, Meier JS, Sapir-Hen L, Stiner MC, Yeshurun R (2018) The emergence of animal management in the southern levant. *Sci Rep* 8(1):9279
- Muthukrishna M, Henrich J (2016) Innovation in the collective brain. *Philosophic Trans Royal Soc B: Biol Sci* 371(1690)
- Nishida T, Uehara S, Nyundo R (1979) Predatory behavior among wild chimpanzees of the mahale mountains. *Primates* 20(1):1–20
- Norris V, Raine DJ (1998) A fission-fusion origin for life. *Origins Life Evol Biosphere: J Int Soc Study Origin Life* 28(4–6):523–537

- Okasha S (2006) Evolution and the levels of selection. Oxford University Press, Oxford
- Okasha S (2007) Multilevel selection and the major transitions in evolution. *Philosophy of Science* 72(5):1013–1025
- Pigliucci M, Müller GB (eds) (2010) Evolution, the extended synthesis. MIT Press, Cambridge
- Pobiner BL (2020) The zooarchaeology and paleoecology of early hominin scavenging. *Evol Anthropol* 29(2):68–82
- Polanyi M (1967) Personal Knowledge: Towards a Post-Critical Philosophy. Horper Torchbooks, New York
- Potts R (1998) Variability selection in hominid evolution. *Evol Anthropol* 7(3):81–96
- Potts R (2012) Environmental and behavioral evidence pertaining to the evolution of early homo. *Curr Anthropol* 53(S6):S299–317
- Premo LS, Tostevin GB (2016) Cultural transmission on the taskscape: exploring the effects of taskscape visibility on cultural diversity. *PLoS One* 11(9):1–22
- Pusey AE (1979) Intercommunity transfer of chimpanzees in Gombe National Park. In: Hamburg DA, McCown ER (eds) The great apes. Benjamin/Cummings, Menlo Park, pp 464–479
- Queller DC (1997) Cooperators since life began. *Q Rev Biol* 72(2):184–188
- Queller DC (2000) Relatedness and the fraternal major transitions. *Philosophic Trans Royal Soc London. Series B, Biol Sci* 355(1403):1647–55
- Queller DC, Strassmann JE (2009) Beyond society: the evolution of organismality. *Philosophic Trans Royal Soc B: Biol Sci* 364(1533):3143–3155
- Querbes A, Vaesen K, Houkes W (2014) Complexity and demographic explanations of cumulative culture. *PloS One* 9(7):e102543
- Radzvilavicius AL, Blackstone NW (2018) The evolution of individuality revisited. *Biol Rev* 93(3):1620–1633
- Read DW (2012) How culture makes us human: primate social evolution and the formation of human societies. Left Coast Press, New York
- Read DW (2020) From *Pan* to *Homo sapiens*: evolution from individual based to group based forms of social cognition, vol 19. Springer, Berlin/Heidelberg
- Read DW, Andersson C (2019) Cultural complexity and complexity evolution. *Adapt Behav* 105971231882229.
- Reindl E, Bandini E, Tennie C (2018) The latent solutions and its relation to the classics: Vygotsky and Köhler. In: Di Paolo L, Di Vincenzo F, De Petrillo F (eds) Evolution of primate social cognition. Springer, Cham, pp 231–248
- Richerson PJ, Baldini R, Bell AV, Demps K, Frost K, Hillis V et al (2016) Cultural group selection plays an essential role in explaining human cooperation: a sketch of the evidence. *Behav Brain Sci* 39:e30
- Richerson PJ, Boyd R, Bettinger RL (2001) Was agriculture impossible during the pleistocene but mandatory during the holocene? A climate change hypothesis. *Am Antiq* 66(3):387–411
- Roche H, Blumenshine RJ, Shea JJ (2009) Origins and adaptations of early *Homo*: what archeology tells us. In: Grine FE, Fleagle JG, Leakey RE (eds) The first humans—origin and early evolution of the genus *Homo*. Springer, Dordrecht, pp 135–147
- Roughley N, Bayertz K (eds) (2019) The normative animal? Oxford University Press, Oxford
- Sandén BA, Hillman KM (2011) A framework for analysis of multi-mode interaction among technologies with examples from the history of alternative transport fuels in Sweden. *Res Policy* 40(3):403–414
- Sanz CM, Morgan DB (2007) Chimpanzee tool technology in the goulougo triangle, Republic of Congo. *J Hum Evol* 52:420–433
- Sayers K, Owen Lovejoy C (2008) The chimpanzee has no clothes. *Curr Anthropol* 49(1):87–114
- Schel AM, Rawlings B, Claidière N, Wilke C, Wathan J, Richardson Jo, Pearson S, Herrelko ES, Whiten A, Slocombe K (2013) Network analysis of social changes in a captive chimpanzee community following the successful integration of two adult groups. *Am J Primatol* 75(3):254–266
- Schiffer MB (2005) The devil is in the details: the cascade model of invention processes. *Am Antiq* 70(3):485–502
- Schöning C, Humle T, Möbius Y, McGrew WC (2008) The nature of culture: technological variation in chimpanzee predation on army ants revisited. *J Hum Evol* 55(1):48–59
- Shea N (2009) Imitation as an inheritance system. *Philosophic Trans Royal Soc London. Series B, Biol Sci* 364(1528):2429–43
- Shea N, Boldt A, Bang D, Yeung N, Heyes C, Frith CD (2014) Supra-personal cognitive control and metacognition. *Trends Cogn Sci* 18(4):186–193
- Shelton DE, Michod RE (2014) Group selection and group adaptation during a major evolutionary transition: insights from the evolution of multicellularity in the volvocine algae. *Biol Theory* 9(4):452–469
- Shelton DE, Michod RE (2020) Group and individual selection during evolutionary transitions in individuality: meanings and partitions. *Proc Royal Soc B* 20190364
- Sherwood CC, Gómez-Robles A (2017) Brain plasticity and human evolution. *Annu Rev Anthropol* 46:399–419
- Sherwood CC, Subiaul F, Zawidzki TW (2008) A natural history of the human mind: tracing evolutionary changes in brain and cognition. *J Anat* 212(4):426–454
- Simon HA (1962) The architecture of complexity. *Proc Am Philos Soc* 6(106):467–482
- Smaldino PE (2014) The cultural evolution of emergent group-level traits. *Behav Brain Sci* 37(3):243–254
- Smith BD (2007) The ultimate ecosystem engineers. *Science* 315(5820):1797–1798
- Smith CM, Gabora L, Gardner-O’Kearny W (2018) The extended evolutionary synthesis facilitates evolutionary models of culture change. *Clodynamics: J Quant History Cult Evol* 9(2)
- Sober E, Wilson DS (1994) A critical review of philosophical work on the units of selection problem. *Philosophy Sci* 61:534–555
- Stankiewicz R (2000) The concept of ‘design space’. In: Ziman J (ed) Technological innovation as an evolutionary process. Cambridge University Press, Cambridge, pp 234–247
- Sterelny K (2011) Evolvability reconsidered. In: Calcott B, Sterelny K (eds) The major transitions in evolution revisited. MIT Press, Cambridge, pp 83–100
- Stiner MC (2002) Carnivory, coevolution, and the geographic spread of the genus homo. *J Archaeol Res* 10(1):1–63
- Stiner MC, Buitenhuis H, Duru G, Kuhn SL, Mentzer SM, Natalie DM et al (2014) A forager-herder trade-off, from broad-spectrum hunting to sheep management at Aşıklı Höyük, Turkey. *Proc Natl Acad Sci USA* 111(23):8404–8409
- Stiner MC, Özbaşaran M, Duru G (2021) Aşıklı Höyük : the generative evolution of a central anatolian PPN settlement in regional context. *J Archaeol Res*
- Stout D, Rogers MJ, Jaeggi AV, Semaw S (2019) Archaeology and the origins of human cumulative culture: a case study from the Earliest Oldowan at Gona, Ethiopia. *Curr Anthropol* 60(3):309–340
- Stout D, Semaw S, Rogers MJ, Cauche D (2010) Technological variation in the earliest Oldowan from Gona, Afar, Ethiopia. *J Hum Evol* 58(6):474–491
- Strassmann JE, Queller DC (2010) The social organism: congresses, parties, and committees. *Evolution* 64(3):605–616
- Szathmáry E (1986) The eukaryotic cell as an information integrator. *Endocytobiosis and Cell Research* 3(2):113–132
- Szathmáry E (2015) Toward major evolutionary transitions theory 2.0. *Proc Natl Acad Sci USA* 112(33):10104–10111

- Szathmáry E, Demeter L (1987) Group selection of early replicators and the origin of life. *J Theor Biol* 128(4):463–486
- Szathmáry E, Maynard-Smith J (1995) The major evolutionary transitions. *Nature* 374(6519):227–232
- Tehrani JJ, Riede F (2008) Towards an archaeology of pedagogy: learning, teaching and the generation of material culture traditions. *World Archaeol* 40(3):316–331
- Tennie C, Call J, Tomasello M (2009) Ratcheting up the ratchet: on the evolution of cumulative culture. *Philosophic Trans Royal Soc London. Series B, Biol Sci* 364(1528):2405–15
- Tennie C, Premo LS, Braun DR, McPherron SP (2017) Early stone tools and cultural transmission: resetting the null hypothesis. *Curr Anthropol* 58(5):000–000
- Theiner G (2014) Varieties of group cognition. *The Routledge Handbook of Embodied Cognition* 362:347–358
- Thompson JL, Nelson AJ (2011) Middle childhood and modern human origins. *Hum Nat* 22(3):249–280
- Thompson JL, Nelson AJ (2016) Childhood and patterns of growth in the genus *Homo*. In: Meehan CL, Crittenden AN (eds) *Childhood: origins, evolution, and implications*. University of New Mexico Press, Albuquerque, pp 75–102
- Thompson JC, Carvalho S, Marean CW, Alemseged Z (2019) Origins of the human predatory pattern: the transition to large-animal exploitation by early hominins. *Curr Anthropol* 60(1):1–23
- Tomasello M (1996) Do apes ape? In: Heyes CM, Galef BG (eds) *Social learning in animals: the roots of culture*. Academic Press, San Diego, pp 319–346
- Tomasello M, Davis-Dasilva M, Camak L, Bard K (1987) Observational learning of tool use by young chimpanzees and enculturated chimpanzees. *Hum Evol* 2(1982):175–183
- Tostevin GB (2007) Social intimacy, artefact visibility and acculturation models of Neanderthal – modern human interaction. In: Mellars PA (ed) *Rethinking the human revolution: new behavioural and biological perspectives on the origin and dispersal of modern humans*. McDonald Institute for Archaeological Research, Cambridge, pp 341–357
- Tostevin GB (2019) Content matters: the materiality of cultural transmission and the intersection of paleolithic archaeology with cultural evolutionary theory. In: Love A, Wimsatt WC (eds) *Beyond the meme: development and structure in cultural evolution*. University of Minnesota Press, Minneapolis, pp 311–364
- Traulsen A, Nowak MA (2006) Evolution of cooperation by multilevel selection. *Proc Natl Acad Sci USA* 103(29):10952–10955
- Ungar PS, Grine FE, Teaford MF (2006) Diet in early homo : a review of the evidence and a new model of adaptive versatility. *Annu Rev Anthropol* 35(1):209–228
- van de Waal E (2018) On the neglected behavioural variation among neighbouring primate groups. *Ethology* 124(12):845–854
- van de Waal E, Borgeaud C, Whiten A (2013) Potent social learning and conformity shape a wild primate's foraging decisions. *Science* 340(6131):483–485
- van Leeuwen EJC, Cronin KA, Daniel BM, Haun RM, Bodamer MD (2012) Neighbouring chimpanzee communities show different preferences in social grooming behaviour. *Proc Royal Soc b: Biol Sci* 279(1746):4362–4367
- van Schaik CP (2016) *The primate origins of human nature*. Wiley, Hoboken
- van de Waal E, Renevey N, Favre CM, Bshary R (2010) Selective attention to philopatric models causes directed social learning in wild vervet monkeys. *Proceed Royal Soc B: Biol Sci* 277(1691):2105–2111
- Watson SK, Reamer LA, Mareno MC, Vale G, Harrison RA, Lambeth SP et al (2017) Socially transmitted diffusion of a novel behavior from subordinate chimpanzees. *Am J Primatol* 79(6):1–10
- West-Eberhard MJ (2003) *Developmental plasticity and evolution*. Oxford University Press, Oxford
- West SA, Toby Kiers E (2009) Evolution: what is an organism? *Curr Biol* 19(23):R1080–R1082
- Whiten A (2005) The second inheritance system of chimpanzees and humans. *Nature* 437(7055):52–55
- Whiten A, Erdal D (2012) the human socio-cognitive niche and its evolutionary origins. *Philosophic Trans Royal Soc B: Biol Sci* 367(1599):2119–2129
- Whiten A, Goodall J, McGrew WC, Toshisada N, Reynoldsk V, Sugiyama Y et al (1999) Cultures in chimpanzees. *Nature* 399:15–18
- Whiten A, Goodall J, McGrew WC, Toshisada N, Reynoldsk V, Sugiyama Y et al (2001) Charting cultural variation in chimpanzees. *Behaviour* 138(11):1481–1516
- Whiten A, Horner V, Marshall-Pescini S (2003) Cultural panthropology. *Evol Anthropol* 12(2):92–105
- Whiten A, Horner V, de Waal FBM (2005) Conformity to cultural norms of tool use in chimpanzees. *Nature* 437(7059):737–740
- Whiten A, McGuigan N, Marshall-Pescini S, Hopper LM (2009) Emulation, imitation, over-imitation and the scope of culture for child and chimpanzee. *Philosophic Trans Royal Soc London. Series B, Biol Sci* 364(1528):2417–28
- Williams GC (1966) *Adaptation and natural selection*. Oxford University Press, Oxford
- Wilson ML (2013) Chimpanzees, warfare, and the invention of peace. In: Fry DP (ed) *War, peace, and human nature: the convergence of evolutionary and cultural views*. Oxford University Press, Oxford, pp 361–388
- Wilson ML, Wrangham RW (2003) Intergroup relations in chimpanzees. *Annu Rev Anthropol* 32(1):363–392
- Wimsatt WC (1975) Complexity and organization. In: *Topics in the philosophy of biology*. Reidel, Dordrecht, pp 174–193
- Wimsatt WC, Griesemer JR (2007) Reproducing entrenchments to scaffold culture: the central role of development in cultural evolution. In: Sansom R, Brandon RN (eds) *Integrating evolution and development: from theory to practice*. MIT Press, Cambridge, MA, pp 227–323
- Wrangham RW (1979) On the evolution of ape social systems. *Soc Sci Inf* 18(3):336–368
- Zeder MA 2012 Pathways to animal domestication. In: Gepts P, Famula TR, Bettinger RL, Brush SB, Damania AB, McGuire PE, Qualset CO (eds) *Biodiversity in agriculture: domestication, evolution and sustainability*. Cambridge University Press, New York, pp 227–259
- Zeder MA (2015) Core questions in domestication research. *Proc Natl Acad Sci USA* 112(11):3191–3198
- Zeder MA (2018) Why evolutionary biology needs anthropology: evaluating core assumptions of the extended evolutionary synthesis. *Evol Anthropol: Issues News Rev* 27(6):267–284
- Ziman J (2000) *Technological innovation as an evolutionary process*. Cambridge University Press, Cambridge

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